



On the role of mechanical feedback in plant morphogenesis

Hadrien Oliveri

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**THÈSE POUR OBTENIR LE GRADE DE DOCTEUR
DE L'UNIVERSITE DE MONTPELLIER**

En Informatique

École doctorale : Information, Structures, Systèmes

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**On the role of mechanical feedback in plant
morphogenesis**

Présentée par Hadrien OLIVERI

Le 28 mai 2019

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**UNIVERSITÉ
DE MONTPELLIER**

To my parents.

La filosofia è scritta in questo grandissimo libro, che continuamente ci sta aperto innanzi agli occhi (io dico l'Universo), ma non si può intendere, se prima non il sapere a intender la lingua, e conoscer i caratteri ne quali è scritto. Egli è scritto in lingua matematica, e i caratteri son triangoli, cerchi ed altre figure geometriche, senza i quali mezzi è impossibile intenderne umanamente parola; senza questi è un aggirarsi vanamente per un oscuro labirinto.

- Galileo Galilei, *Il Saggiatore*, Rome, 1623

Abstract

How do living objects acquire their shape? Incontrovertibly, *morphogenesis* is largely regulated by genes. Yet, the precise link between the chemical processes associated with genes, on the one hand, and geometry, on the other hand, is not completely identified. This link is most probably indirect, and mediated by mechanical processes. It is now well accepted that intracellular molecular processes regulate locally cell mechanical properties and that shape emerges as the global resolution of resulting mechanical constraints.

This so-called biomechanical paradigm is employed in this thesis in the context of plant morphogenesis, that mostly relies on cell growth. The local control of growth is crucial for the stability and robustness of morphogenesis, and relies on various regulatory mechanisms. In particular, according to a recent hypothesis, cells may dynamically adapt their growth behavior in response to the mechanical forces they experience.

This local regulation integrates at larger, multicellular scale, in a nonintuitive way. In this thesis, I investigate i/ the mathematical formalization of a stress-based control of growth and ii/ the macroscopic emergent behavior of such mechanism. To do so, I have used a multiscale modeling approach, based on a continuum mathematical model of growth (previously developed within the theory of *morphoelasticity*), and on a mean description of the molecular processes supposedly involved in mechanoperception and the control of cell elastic properties. To study this model, I have designed dedicated algorithms, integrated into a previously developed software environment, based on the *finite element method*. This model is then used to study the mechanical stability of highly asymmetric organs like leaves, suggesting that a force-based control of growth allows for the amplification of shape asymmetry during development.

Résumé

L'acquisition de la forme – ou *morphogenèse* – chez les systèmes vivants, est largement contrôlée par les gènes. Néanmoins, le lien précis entre, d'une part, les processus chimiques locaux associés aux gènes, et, d'autre part, la géométrie des tissus, n'est pas complètement identifié. Ce lien est vraisemblablement très indirect et médié par des processus mécaniques. Ainsi, il est aujourd'hui admis que les processus chimiques intracellulaires régulent les propriétés mécaniques des cellules seulement localement, et que la forme émerge comme la résolution globale de contraintes mécaniques.

Ce paradigme, dit *biomécanique*, est employé dans cette thèse dans le cas de la morphogenèse des plantes, qui repose majoritairement sur la croissance cellulaire. Le contrôle local de cette croissance est crucial pour la stabilité et la robustesse de la morphogenèse, et implique différents mécanismes de régulation. En particulier, selon une hypothèse récente, les cellules pourraient adapter dynamiquement leur croissance en réponse aux forces qu'elles subissent.

Cette régulation locale s'intègre à une échelle multicellulaire de manière non intuitive. Dans cette thèse, j'ai exploré i/ une formalisation mathématique de la régulation de la croissance par les contraintes mécaniques et ii/ le comportement macroscopique émergent d'un tel mécanisme. Pour cela, j'ai adopté une approche de modélisation multi-échelle basée sur une formulation mathématique continue de la croissance cellulaire (développée précédemment dans le cadre de la théorie de la *morphoélasticité*), et sur une description moyenne des processus moléculaires locaux étant supposés impliqués dans la *mécano-perception* et le contrôle de l'élasticité des cellules. J'ai d'autre part conçu des algorithmes dédiés à l'étude de ce modèle, intégrés dans un environnement logiciel existant, basé sur la *méthode des éléments finis*. Ce modèle est en particulier utilisé dans l'étude de la stabilité d'organes à fort degré d'asymétrie, tels que les feuilles, en suggérant qu'un contrôle de la croissance basé sur les forces peut permettre l'amplification d'asymétries initiales.

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Chapter 0

Introduction

0.1 General context: modeling developmental biology

How living objects acquire their shape has intrigued inquisitive minds for centuries and is still an open question (see [Coen, 2000](#), for a layman's historical introduction to developmental biology). Incontrovertibly, this crucial aspect of biological development (known as *morphogenesis*) is for great part regulated by genes. Within a tissue, those may be expressed non-homogeneously, instructing each cell, more or less directly, with a specific behavior. For this reason, the study of genes and their translation into proteins is a cornerstone of developmental biology, which has seen increasing scientific and technological investments in the last decades.

However, living things are often *more than the sum of their parts*: knowing the individual behavior of each individual cell (or molecule) does not completely inform us on the whole body behavior. This remarkable feature of complex – in particular living – systems (known as *emergence*) was well illustrated by various attempts to simulate life *in silico*, for example in cellular automata like Conway's *Game of Life* ([Berlekamp et al., 2004](#)) or Langton's ant ([Langton, 1986](#)). Relying on minimal local rules governing their temporal evolution, these artificial systems may display rich and highly unpredictable large-scale behaviors.

Genetic processes undoubtedly play a central role in providing cells with *properties* (their local "rules"). However, they probably do not directly hard code the global behavior of a developing organism, which mostly emerges from more or less local cell-cell interactions. In particular the role of genes on the establishment of global shape is most plausibly indirect, and our understanding of it remains fragmented.

Without denying the crucial role of biochemistry, D'Arcy Thompson, in his pioneering book entitled *On growth and form* (Thompson, 1917), stressed that morphogenesis is by essence a physical and mechanical process (see also Hamant, 2017, for an interesting appreciation of Thompson's work). Mechanics is by definition concerned with the motion and equilibria of physical bodies, and living things do not escape its constraints. The so-called *biomechanical* approach aims at explaining biological behaviors by seeing them as realizations of more general mechanical and physical principles. In this view, it is commonly believed that mechanics is the ultimate link between genes and shape.

Mathematical modeling provides the essential basis to describe mechanics, by formalizing notions like energy, deformation and force, in a quantitative manner. Yet, often enough, the complex synergy between mechanical effects, genetics and other physical phenomena, escapes a complete analytic mathematical understanding. In such a case, we may resort to numerical computational models. These allow to systematically take into account numerous hypotheses, scales and actors, as well as the couplings that may exist between them over time, beyond qualitative intuition.

0.2 Scope of the thesis

In this work, I study the dynamic control of plant morphogenesis. Plant shape depends on the individual growth of the cells composing a *tissue*. How cells collectively achieve cohesive growth and globally repeatable morphogenesis is a highly debated question, that involves studying local controls and integrated large scale effects. These control may depend on genes and on numerous other biophysical processes that, combined together, may produce various feedback loops. One of these, recently proposed, may integrate mechanics itself as a central cue of growth control (Hamant et al., 2008). According to this hypothesis, cells would in fact dynamically adapt their growth behavior according to the mechanical forces that they experience. The core question of this thesis is to understand what collective behavior can emerge from such a control and how morphogenesis can be regulated through it.

To address this question, we developed a biomechanical approach. Plant cell growth is fueled by important intracellular turgor-induced forces (in plants, *turgor pressure* can build up to several atmospheres, Hüsken et al., 1978). Under this loading, plant cells deform, in a way that depends in particular on the time scale of the observation. At short time scale, they display solid, elastic mechanical properties, while, at longer time scale they may deform in a plastic (irreversible) manner, which defines cell growth.

This kind of nontrivial process, in general, is the object of the theory of *morphoelasticity* (exposed later on), that aims at mathematically characterizing the link between the shape and the mechanics of biological objects. This theory is in particular based on a two time scale resolution of growth and on a separation between the elastic and plastic components of deformation. In this thesis, I use a computational approach based on these general principles to model the force-based control of plant tissue growth.

Chapter 1

State of the art

1.1 Morphogenesis at the *shoot apical meristem*

Unlike animals, most plants continuously generate organs throughout their lifespan. In higher plants, the formation of those organs (shoots, roots, flowers, leaves, etc.), termed *organogenesis*, occurs at different precise zones of the plant named *meristems*.

The aboveground organs (e.g. leaves, flowers) in particular, are formed at the *shoot apical meristem (SAM)*¹ (Fig. 1.1(a)). The SAM is a dome-shaped pool of stem cells, located at the tip of the plant axis (Ha et al., 2010, Steeves and Sussex, 1989, Traas and Doonan, 2001). It produces stem tissues (resulting in the axial expansion of the shoot) and initiates lateral organs (branches, leaves, flowers). Organ initiation occurs periodically, at the periphery of the meristem's *central zone*. As organs are initiated and expand, the meristem tip progresses due to the axial growth of the meristem, making room for next organs to appear (Fig. 1.1(b)).

Organogenesis at the SAM is spatially and temporally regular (following a given *phylloclastic* spatiotemporal program, Traas, 2013) and organ shape is highly conserved. This yet relies on the collective behavior of many cells that, individually, have only a local "knowledge" of the global picture. To solve this apparent paradox, a high level of regulation involving some type of cell-cell communication is necessary, guaranteeing a globally cohesive cell behavior. Supporting this view, several mechanisms have been identified as crucial regulators of morphogenesis – e.g. auxin transport (see Sassi and Vernoux, 2013, for a review), architect genes (Weigel and Meyerowitz, 1994) or mechanosensing (Hamant et al., 2008) – without which the regularity of organogenesis

¹There are other types of meristems (*root meristems, axillary meristems, cambium, etc.*).

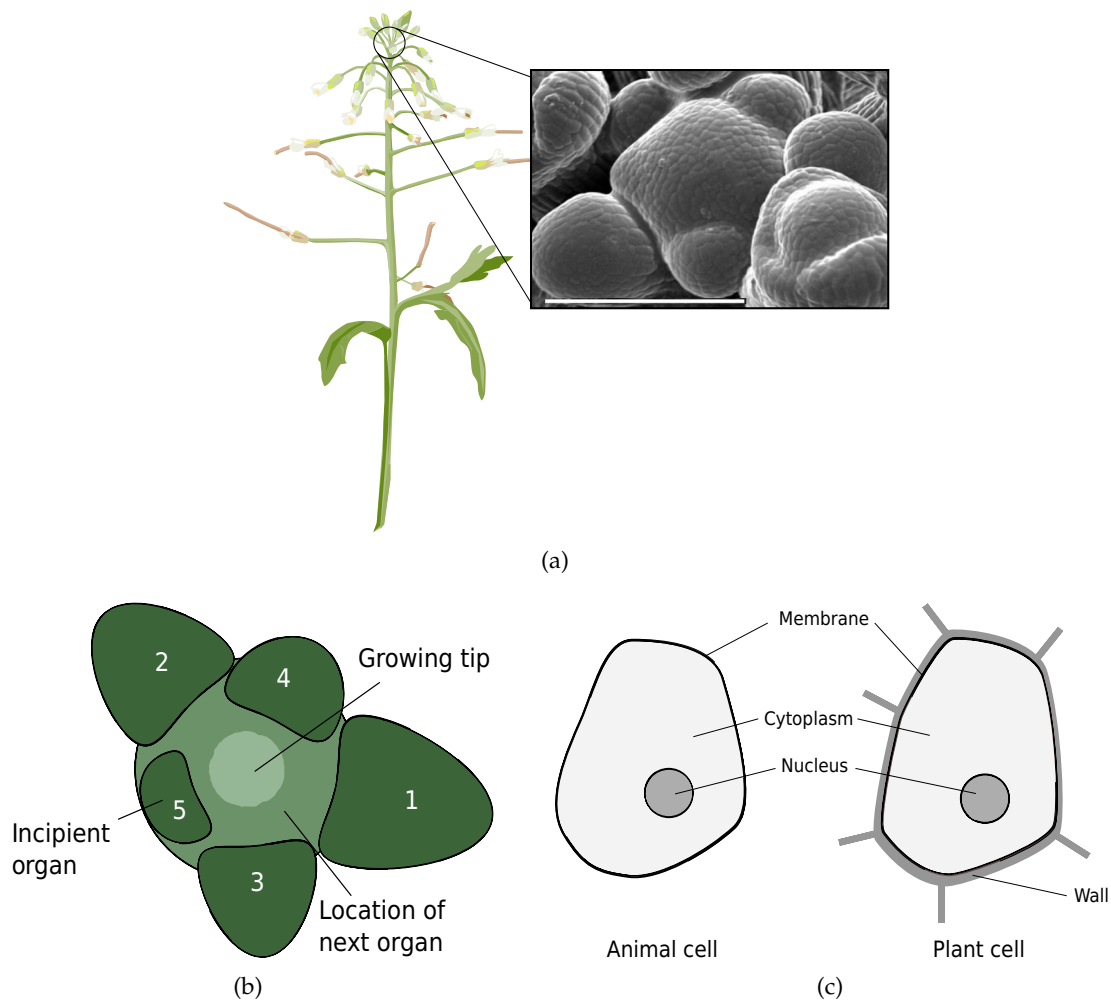


Figure 1.1: (a) Cartoon of *Arabidopsis thaliana*'s inflorescence (Image: www.pflanzenforschung.de) and close-up on the inflorescence meristem (scanning electron microscopy, scale bar = 100 μm , Image courtesy of J. Traas). (b) General structure of the shoot apical meristem (top view). Organs form at the periphery of the meristem at a regular pace, as the meristem tip progresses. (c) General structure of the animal and plant cells. Unlike animal cells, plant cells are embedded in a stiff wall. This cell wall prevents relative cell displacements, as two cells cannot detach from each other. By contrast, animal cells, that have their membranes in contact, may slide against one another more freely. Redrawn from Coen (2000).

is dramatically impaired. To provide a rational, integrated view of how these mechanisms control morphogenesis in time and space, mathematical and computational models have been designed, in the form of virtual tissues coupling various biological features (Chickarmane et al., 2010). In particular, one essential challenge in constructing these models is how to take growth into account.

In plants, cells are embedded within a stiff and sticky exoskeleton, the *cell wall*, that prevents them from moving relative to one another (Fig. 1.1(c)). In walled cells, *osmosis* results in a balance between cell *turgor pressure* and cell wall elastic stretch. Under

sufficient turgor-induced tension, the cell wall may expand irreversibly, giving the cell room to absorb more water and augment its volume (see [Ali et al., 2014](#), [Hamant and Traas, 2010](#), for reviews). This irreversible process defines *cell growth*. In a multicellular context, due to cell-cell, short-range interactions, pressure forces not only affect cells autonomously, but also propagate across the tissue (see Fig. 1.2 and [Boudon et al., 2015](#)). Local growth then results from the integration of both long and short range mechanical influences. To understand how those mechanical interactions integrate at the scale of an entire tissue, a number of biomechanical approaches have been proposed..

In the sequel, I critically review the various models published. I first present a kinematic description of plant growth, *i.e.* the set of variables that allow to describe the motion, independent of its causes, and second, an overview of the kinetics of growth – *i.e.* the link between motions/equilibria and forces – that in plants has been described in various manners.

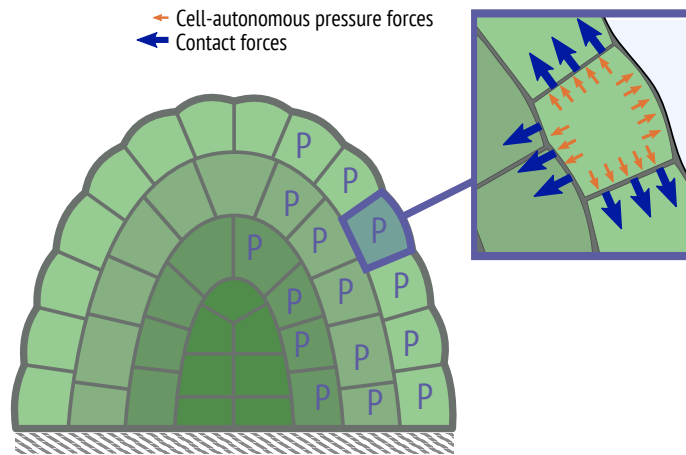


Figure 1.2: Within a multicellular context, the wall is subject to both cell-autonomous pressure-induced forces and cell-cell contact forces.

1.2 Kinematics of growth

1.2.1 Kinematic parameters of tissue motion

Meristem development excludes cell migration and *apoptosis* (*i.e.* programmed cell death)². In this context, topological changes in the tissue result exclusively from *cell divisions*, and organ shape changes can only emerge from individual cell dilation. It was then proposed that complex morphogenesis can be achieved through spatial heterogeneities and anisotropies in the individual cell dilation speeds, *i.e.* through the fact

²By contrast, in animals, cells are freer to slip against one another. Many animal developmental processes involve collective cell migration ([Friedl and Gilmour, 2009](#)) or apoptosis (e.g. formation of digits, [Jacobson et al., 1997](#)).

that some cells (or groups of cells) grow faster than their neighbors and with a rate that depends on direction (Coen et al., 2004).

Authors have stressed that, although overall shape deformation may be complex, it can be decomposed into a limited set of *kinematic* parameters (i.e. descriptors of the motion, independent of its causes), if taken at a sufficiently small scale (Avery, 1933, Coen et al., 2004, Erickson, 1976). Those parameters are the main *direction* of the deformation, its degree of *anisotropy*, its *volumetric expansion rate* and its *rotation rate* (Coen et al., 2004).

Mathematically, this can be formalized in the framework of continuum kinematics (see for instance Holzapfel, 2000, for an introduction). Assume that at a given time t , each point \mathbf{x} of the tissue continuum moves relative to the lab frame with an instantaneous velocity $\mathbf{v}(\mathbf{x}, t)$ (Fig. 1.3(a)). Clearly, $\mathbf{v}(\mathbf{x}, t)$ gives little information on the growth at \mathbf{x} , since it may encompass all kinds of motion, including, in particular, rigid transformations of the whole domain. A more meaningful descriptor of shape change would focus on the relative – rather than absolute – motion of the domain points. This relative motion is quantified by the *velocity gradient* (often noted \mathbf{L}), that compares the instantaneous velocity at the control point \mathbf{x} to that at neighboring positions:

$$\mathbf{L}(\mathbf{x}, t) \stackrel{\text{def.}}{=} \frac{\partial \mathbf{v}}{\partial \mathbf{x}}(\mathbf{x}, t) \quad \left(\text{i.e. } L_{ij} = \frac{\partial v_i}{\partial x_j}, \quad i, j = 1, 2, 3 \right). \quad (1.1)$$

The velocity gradient \mathbf{L} is a second order tensor field that describes the instantaneous linear transformation that affects an infinitesimal volume at the vicinity of point \mathbf{x} . In 1D, the velocity gradient describes the instantaneous lengthening/shortening of a given length element (see Fig. 1.3(b)). In general, this transformation may affect the size and shape of this volume, and its orientation (while differentiation eliminates uniform components, i.e. translations). This can be evidenced by splitting \mathbf{L} into its symmetric ($\mathbf{\Sigma}$) and skew-symmetric ($\mathbf{\Omega}$) parts:

$$\mathbf{L} = \mathbf{\Sigma} + \mathbf{\Omega}, \quad \text{with} \quad \mathbf{\Sigma} \stackrel{\text{def.}}{=} \frac{1}{2} (\mathbf{L} + \mathbf{L}^T) \quad \text{and} \quad \mathbf{\Omega} \stackrel{\text{def.}}{=} \frac{1}{2} (\mathbf{L} - \mathbf{L}^T), \quad (1.2)$$

respectively termed *Eulerian strain rate tensor* and *spin tensor* (Fig. 1.3(c)).

As $\mathbf{\Sigma}$ is by construction a symmetric tensor (and positive since growth is an expansion process), it defines a direction of main extension (its first *eigenspace*), a dilation amplitude (its trace) and three degrees of dilation anisotropy (e.g. the pairwise ratios between the three eigenvalues). These anisotropies reduce to a single one, if we restrict ourselves to observe deformations in two dimensions. Note that tensor $\mathbf{\Sigma}$ vanishes in the case of a rigid motion.

As for the spin tensor Ω , it quantifies the rotational motion of particles and is dual to the *instantaneous angular velocity vector* ω :

$$\Omega \cdot \mathbf{h} = \omega \times \mathbf{h} \quad \text{where} \quad 2\omega = \text{curl} \mathbf{v} \quad (1.3)$$

for any vector \mathbf{h} (operator ' \times ' depicts the vector cross product). Equation (1.3) makes the link between Ω and rotation speed more apparent. Further details may be found in [Holzapfel \(2000\)](#) (see chapters *Algebra of tensors* and *Rates of deformation tensors*).

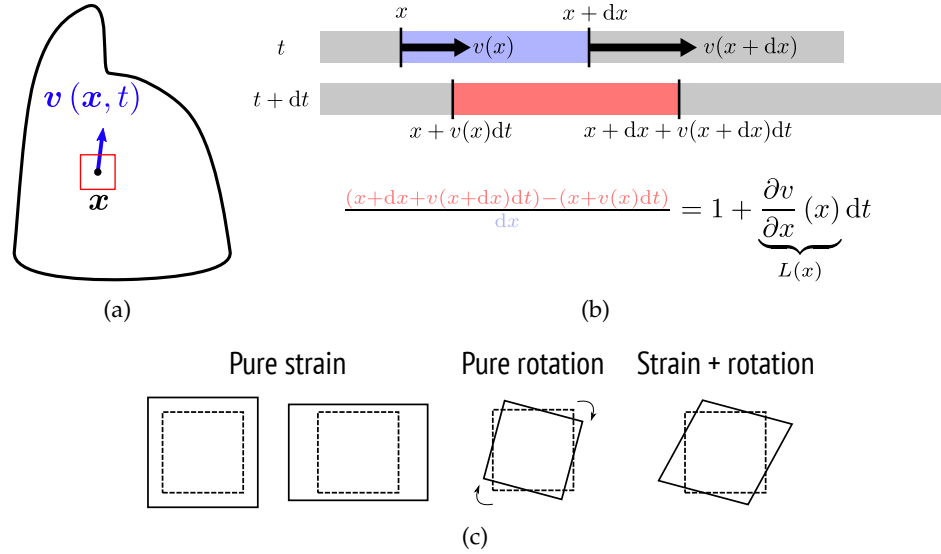


Figure 1.3: (a) Instantaneous velocity at position x . (b) Growth of a 1D rod. Between t and $t + dt$, the infinitesimal portion of length dx as moved and increased in length. The new length is obtained by subtracting the position of its right extremity $x + dx + v(x + dx)dt$ and that of its left extremity $x + v(x)dt$. The line dilation rate of the element is equal to the ratio of the new length to the previous length, or $1 + L(x)dt$. (c) Schematic representations of the various kind of deformations applied to an infinitesimal square volume. Pure strain, pure rotation, and general deformation.

1.2.2 General principles of morphoelasticity

To model the physics of biological growth, authors stressed that part of the observed deformation is *reversible* and governed through elastic processes, and the other part is *irreversible*, governed through plastic processes. Mathematically, this idea was expressed in a continuum formalization originally proposed in [Rodriguez et al. \(1994\)](#) and generally referred as *morphoelasticity* (see also [Garikipati, 2009](#)). This approach, summarized hereafter, was subsequently developed in [Goriely and Ben Amar \(2007\)](#) which discusses a general procedure to represent the continuous growth of an elastic domain in the formalism proposed by Rodriguez and colleagues (see also [Goriely and Moulton, 2011](#)).

Let's assume that at initial time t_0 (resp. at time $t \geq t_0$) the tissue is defined by an initial *reference* configuration $\mathcal{B}(t_0)$ (resp. *current* configuration $\mathcal{B}(t)$). We consider the mapping $\mathbf{x} : \mathbf{X} \in \mathcal{B}(t_0) \rightarrow \mathcal{B}(t)$ that associates each material particle of the domain in the configuration $\mathcal{B}(t_0)$ with its position at t in $\mathcal{B}(t)$ (Fig. 1.4(a)). As in Section 1.2, the mapping transforms the infinitesimal volumes of $\mathcal{B}(t_0)$ according to a linear operation defined by the gradient of the mapping, that is termed *deformation gradient* and noted \mathbf{F} :

$$\mathbf{F} \stackrel{\text{def.}}{=} \frac{\partial \mathbf{x}}{\partial \mathbf{X}}. \quad (1.4)$$

Remark that, by virtue of the chain rule³, the previously-introduced velocity gradient \mathbf{L} (Eq. (1.1)) can be related to the time derivative of \mathbf{F} according to:

$$\mathbf{L} = \frac{\partial \mathbf{F}}{\partial t} \cdot \mathbf{F}^{-1}. \quad (1.5)$$

The observed configuration $\mathcal{B}(t)$ generally contains mechanical *stresses* (residual or due to external forces). The fundamental postulate of morphoelasticity – formulated by Rodriguez and coworkers – is that, at any time, it exists a hidden stress-free state of the system, obtained by *virtually* suppressing external loading and “detaching” the tissue pieces apart from each other (Fig. 1.4(b)). By definition, growth is the change of this stress-free state with time, quantified by the *growth tensor* \mathbf{F}_g (the subscript ‘g’ stands for *growth*), that maps the initial (stress-free) configuration $\mathcal{B}(t_0)$ onto the current stress-free configuration $\mathcal{B}'(t)$. Note that there is no requirement for the deformation field \mathbf{F}_g to be compatible, i.e. to derive from a continuous displacement field. Indeed, as the control of growth is supposedly very local, the tissue pieces may readily grow discontinuously, hence no such compatible displacement exists in general. Compatibility is assured by means of the elastic deformation \mathbf{F}_e ⁴ (the subscript ‘e’ stands for *elastic*), at the price of introducing mechanical stresses. The total deformation is the composition of both deformations, which translates into the following *multiplicative decomposition*:

$$\underbrace{\mathbf{F}}_{\mathcal{B}(t_0) \rightarrow \mathcal{B}(t)} = \underbrace{\mathbf{F}_e}_{\mathcal{B}'(t) \rightarrow \mathcal{B}(t)} \cdot \underbrace{\mathbf{F}_g}_{\mathcal{B}(t_0) \rightarrow \mathcal{B}'(t)}. \quad (1.6)$$

We assume that only the elastic contribution \mathbf{F}_e affects the *strain energy density* Ψ of the system:

$$\Psi = \Psi(\mathbf{F}_e). \quad (1.7)$$

³ $\mathbf{L} = \frac{\partial \mathbf{v}}{\partial \mathbf{x}} = \frac{\partial \mathbf{v}}{\partial \mathbf{X}} \cdot \frac{\partial \mathbf{X}}{\partial \mathbf{x}} = \frac{\partial}{\partial \mathbf{X}} \left(\frac{\partial \mathbf{x}}{\partial t} \right) \cdot \frac{\partial \mathbf{X}}{\partial \mathbf{x}} = \frac{\partial}{\partial t} \left(\frac{\partial \mathbf{x}}{\partial \mathbf{X}} \right) \cdot \frac{\partial \mathbf{X}}{\partial \mathbf{x}} = \frac{\partial \mathbf{F}}{\partial t} \cdot \mathbf{F}^{-1}$.

⁴ Again, \mathbf{F}_e does not define a compatible displacement if \mathbf{F}_g does not.

The variation of \mathbf{F}_g may be quantified by the following measure of the *rate of growth*:

$$\mathbf{L}_g \stackrel{\text{def.}}{=} \frac{\partial \mathbf{F}_g}{\partial t} \cdot \mathbf{F}_g^{-1}. \quad (1.8)$$

that quantifies the growth-related part of the total velocity gradient (Figs. 1.4(b) and 1.4(c)).

Note that the decomposition expressed by Eq. (1.6) applies continuously in time (Goriely and Ben Amar, 2007). In the case of biological growth, the time scale associated with the elastic and viscous effects is generally several orders of magnitude smaller than that of the biological effects governing growth. Thereby, the transient viscoelastic regime is generally neglected and the system is considered in a state of quasi-static elastic equilibrium. Based on this assumption, Goriely and Ben Amar have discussed a general algorithmic method to compute growth, consisting of a succession of small incremental elastic relaxations, and small incremental irreversible deformations (see Goriely and Ben Amar, 2007, and Fig. 1.4(b)).

This general view is frequent in the computational modeling of plant growth, although originally, morphoelasticity describes the growth of elastic *continua* specifically (which does not apply to all models). We shall see that plant growth models often feature the most essential properties of morphoelastic materials, that are i/ the separation of irreversible and reversible deformations and ii/ the separation of elastic and plastic time scales. In an effort to uniformize concepts and notations, I will use the previous formalism as a conceptual backbone in what follows.

So far in this chapter, no requirement on the biophysics governing both \mathbf{F}_g and \mathbf{F}_e in time has been given. In particular, \mathbf{F}_g may depend on various quantities, like mechanical quantities (strain or stress), genes, temperature or nutrient concentration for example. In the sequel, I detail how different choices in implementing the elastic and plastic regimes have resulted in different models. I will mostly focus on morphoelastic-type modeling approaches, from a very general standpoint. Accordingly, I chose not to comment agent-based models like Jönsson et al. (2006), cellular Potts models like Grieneisen et al. (2007), or geometric (non mechanical) models like Holloway and Harrison (1999, 2007), Smith et al. (2006). A specificity of the morphoelastic approach is that the energetic state of the system is given by a displacement field, relative to a *reference configuration*, in contrast to viscous approaches (like Cheddadi et al., 2019, Dupuy et al., 2006), that express forces as a function of the *rate* of deformation. For the sake of simplicity and conciseness, some licence will be taken in the treatment of these cited models, as they will not systematically be the object of a separate review.

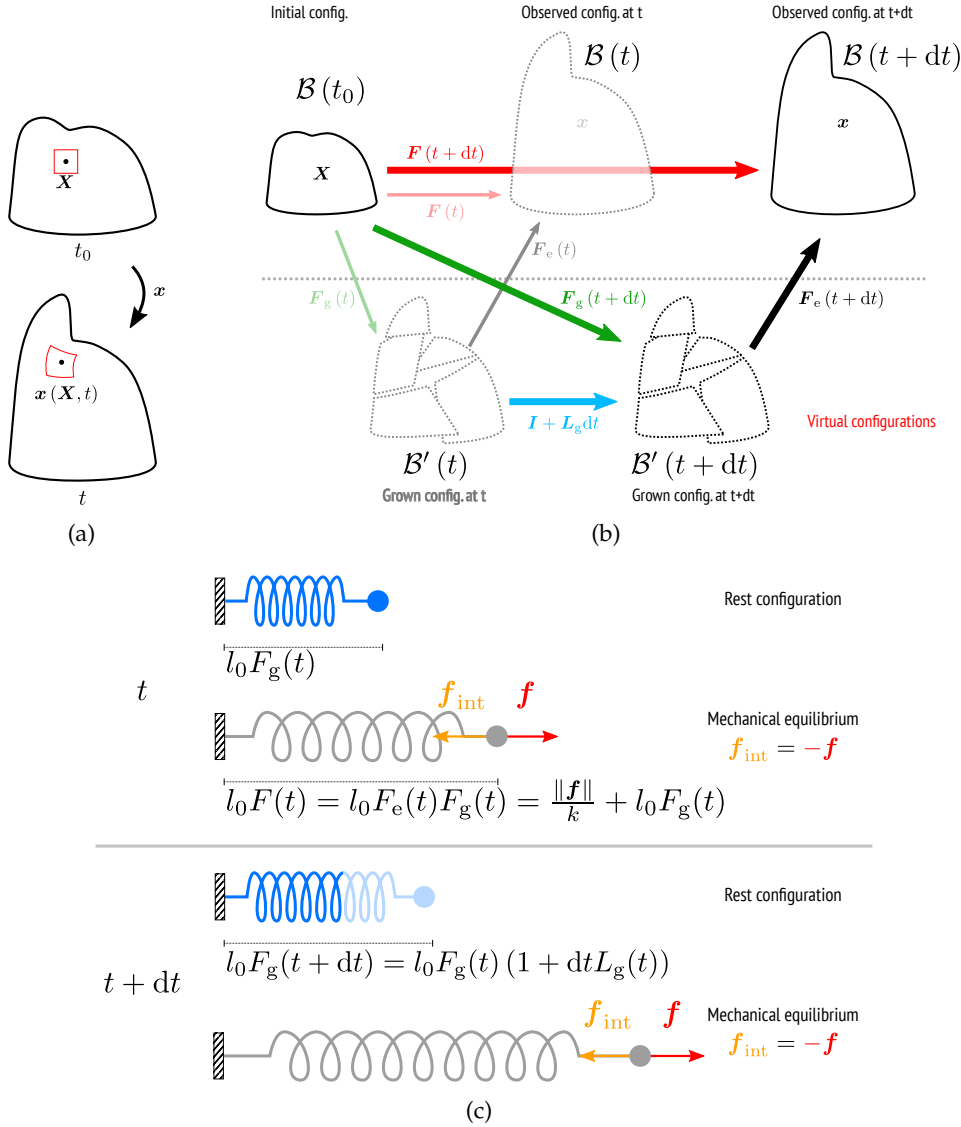


Figure 1.4: (a) Cartoon of the mapping from initial configuration (at t_0) to current configuration (at t). (b) Schematic of the growth of a morphoelastic material. The current configuration is obtained as the combination of both a plastic deformation and an elastic deformation. Growth consists of a modification of the stress-free configuration of the system, that is virtual and does define a compatible displacement field in general. (c) One-dimensional schematic of the growth of a linear spring. At time t the spring has a rest length $l_0 F_g(t)$ (where l_0 depicts the initial rest length), and an observed length $l(t) = l_0 F(t) > l_0(t)$ that results from the loading f (and depends on the spring constant k). The irreversible expansion of the spring is accounted for by an increase of its rest length, that results in a new equilibrium.

1.3 Kinetics of growth

1.3.1 Computing static equilibrium

The notion of mechanical equilibrium may be defined for a conservative system through the *minimum potential energy* principle (see [Holzapfel, 2000](#), [Lanczos, 2012](#), for an introduction to the variational principles of mechanics). In principle, there exists a potential energy functional $\Pi(\mathbf{F}_e)$ for both stress and loads, associated with the system's configuration. The minimum potential energy principle dictates that an equilibrium of the system corresponds to a state for which the total potential $\Pi(\mathbf{F}_e)$ is stationary. If an explicit form of Π can be found, one can consider the problem of finding an equilibrium as an optimization problem.

In a number of cases however, this formalism cannot be employed. This is the case for example when the system is non-conservative, or when the external work depends on the movement of the system in a nontrivial manner (e.g. work of pressure forces when displacement is large). In such cases, a Newtonian, force-based formulation can be employed, consisting of solving a balance of force. In the computational biomechanical models of plants, both energy-based and force-based approaches have been employed.

Several authors have developed 2D force-based vertex models, representing the tissue in the form of a network of 1D mechanical elements such as linear springs ([Bessonov et al., 2013](#), [Cheddadi et al., 2019](#), [Rudge and Haseloff, 2005](#), [Sapala et al., 2018](#)) possibly coupled with torsion springs ([Bessonov et al., 2013](#), [Sapala et al., 2018](#)); or beam finite elements representing both tension and bending ([Dupuy et al., 2010, 2006](#)). In these models, each cell was represented as a polygon composed of discrete mass nodes connected through one-dimensional arcs (Fig. 1.5).

The equilibrium is obtained by balancing the pressure forces with the internal tissue forces. This was done either by integrating the second order equations of motion at each node, until all the node forces fall below a given threshold (for example in [Sapala et al., 2018](#)); or through a first order approximation of the equation of motion, that assumes a strong viscous drag allowing to neglect inertia ([Bessonov et al., 2013](#)).

Energy-based formulations using similar descriptions of the tissue topology were also proposed; in 2D ([Alim et al., 2012](#), [Corson et al., 2009](#), [Louveau et al., 2016](#), [Merks et al., 2011](#), [Qi et al., 2017](#)) or 2D embedded in 3D ([Hamant et al., 2008](#)). In [Corson et al. \(2009\)](#), [Hamant et al. \(2008\)](#), the potential energy was expressed as the sum of the elastic energy (associated with edge deformation) and an external potential explicitly due to pressure. More abstract formulations were chosen in [Alim et al. \(2012\)](#), [Louveau](#)

et al. (2016), Merks et al. (2011), Qi et al. (2017). These rely on an approach often termed *generalized Hamiltonian* formulation, that allows to easily implement specific cell behaviors (e.g. anisotropic growth), e.g. by penalizing deviations from target cell shape (Alim et al., 2012, Louveaux et al., 2016, Merks et al., 2011) or area (Qi et al., 2017). Hamiltonian-based biomechanical modeling generally permits a broad variety of more or less arbitrary energetic contributions, sometimes at the expense of a clear connection with biophysical reality.

From a numerical point of view, Hamiltonian-based methods consider the optimization problem consisting of minimizing the total potential energy Π , that is explicitly formulated. One common heuristic is stochastic optimization, like *Metropolis dynamics* (Metropolis et al., 1953), that is used in Cieslak et al. (2016), Merks et al. (2011), Qi et al. (2017). In a nutshell, this algorithm explores the space of configurations (i.e. the possible moves) in a pseudo-random manner, that consists of looking for energetically more favorable moves, while occasionally authorizing slight increases of energy with a certain probability (in order to escape a non globally-optimal minimum). By contrast, the model detailed in Louveaux et al. (2016) was based on a *quasi-Newton* minimization (the *Broyden-Fletcher-Goldfarb-Shanno* method, see Press et al., 2007) that is deterministic, and as the classical Newton-Raphson minimization method, consists of looking for a root of the energy gradient⁵. To my knowledge, the authors of other cited works (i.e. Alim et al., 2012, Corson et al., 2009, Hamant et al., 2008) did not make mention of the optimization method they used.

In general, vertex-based methods like mass-spring models are relatively easy to implement, benefit from low computational costs, and provide tractable modeling frameworks to represent complex geometries and cell division. Nonetheless, they lack accuracy in approximating some specific properties of continuum multidimensional mechanics, such as shear or incompressibility (Gelder, 1998). In particular, the mechanics of in-plane walls is in general not represented, or at most, in an abstract manner. In addition, these models are restricted to 2D analysis (or 2D embedded in 3D in the only case of Hamant et al., 2008).

To alleviate these limitations, higher-dimensional growth models have been developed. In the line of energy-based approaches like Alim et al. (2012), Corson et al. (2009), Louveaux et al. (2016), Merks et al. (2011), Qi et al. (2017), Cieslak and coworkers have used a generalized Hamiltonian formulation in a 3D model, where the tissue is represented as a collection of connected compartments (Cieslak et al., 2016). In this model, the potential energy stored by each compartment is a function of its volume only. This

⁵In a certain sense, this amounts to looking for a balance of force, yet based on an explicit formulation of the total potential.

is a coarse approximation, and this approach will fail in capturing the elasticity of the tissue more in general. For this purpose, a multidimensional formulation of the strain energy functional is required.

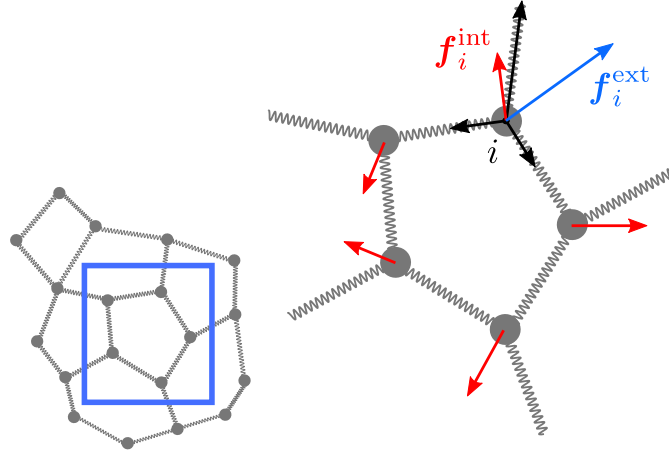


Figure 1.5: Cartoon of a mass-spring system modeling a multicellular tissue.

To provide a more proper physical description of the elastic regime, multidimensional continuum approaches have been proposed to describe tissue elasticity. These generally consider the strain energy density as a function of several variables, depending on multidimensional strain. The *finite element method* (FEM) has become overwhelmingly popular in simulating this kind of system (for an introduction to the general FEM, see for example Hughes (2012), or Zienkiewicz et al. (2000), for applications to solid mechanics). The FEM is a mathematically-sound, generic and efficient method, often used for structural mechanics, and implemented by a myriad of open source or proprietary packages⁶.

In modeling plant morphogenesis, authors have benefited from these convenient properties in modeling tissues in various forms and at different scales, yet with roughly comparable implementations. This was done for example in Fayant et al. (2010), detailing a single-cell model where the cell wall is represented with 2D shell finite elements, which take into account both tangential stress and bending moment. Larger multicellular tissues have also been represented. Fozard and coworkers, for example have extended the previously cited vertex-based models by simulating the mechanics of in-plane walls using planar membrane finite element, that only take into account tangential strains (Fozard et al., 2013). Others have modeled the tissue in 3D, as an assembly of 2D meshed surfaces mimicking the network of cell walls forming the tissue, using 2D shell elements (Robinson and Kuhlemeier, 2018, Sampathkumar et al., 2014a) or simpler membrane elements (Bassel et al., 2014, Boudon et al., 2015, Hamant et al., 2008). Other models have considered larger-scale tissues seen as continuum domains

⁶See for example en.wikipedia.org/wiki/List_of_finite_element_software_packages.

(without cells), in 2D in [Hervieux et al. \(2016\)](#), [Hong et al. \(2016\)](#) (that employ a model originally developed in [Bonazzi et al., 2014](#), for fission yeast) and [Coen et al. \(2004\)](#), [Kuchen et al. \(2012\)](#), 2D embedded in 3D ([Kierzkowski et al., 2012](#)), or 3D ([Kennaway et al., 2011](#)).

A different approach was chosen by Bozorg and colleagues who used the *triangular biquadratic springs* (developed in [Delingette, 2008](#)). Triangular biquadratic springs build a formal equivalence between spring-based modeling and membrane mechanics. This approach was used to model the tissue either as a pressurized membrane ([Bozorg, 2016](#), [Bozorg et al., 2014](#)); or as a multicellular complex ([Daher et al., 2018](#)) similar to that modeled in [Bassel et al. \(2014\)](#), [Boudon et al. \(2015\)](#), [Mosca et al. \(2017\)](#), [Robinson and Kuhlemeier \(2018\)](#).

Generally, the cited authors have considered elements under finite (by contrast with *infinitesimal*) deformations, which is usually associated with a force-based resolution of the equilibrium, computed by solving the second ([Bozorg et al., 2014](#)) or first order ([Bassel et al., 2014](#), [Boudon et al., 2015](#), [Bozorg et al., 2016](#)) equation of motion, through various forward or backward integration procedures. At each integration step, external forces are computed from pressure and current geometry, while (internal) reaction forces are obtained as the negative gradient of the strain energy (itself often expressed based on Saint-Venant Kirchhoff-like hyperelastic formulations, see [Holzapfel, 2000](#)). By contrast, in small deformations (e.g. in [Hervieux et al., 2017](#), [Hong et al., 2016](#), [Kennaway et al., 2011](#)), equilibrium can be directly found using a static FEM resolution, that consists of considering the equilibrium as the solution of a linear system, itself derived from a variational formulation ([Zienkiewicz et al., 2000](#)). When coupled with growth, this last solution generally requires a very small growth time step size to be accurate.

While more or less complex models have been proposed to express the stretch of the wall itself, the loading forces and their source (turgor pressure) were generally represented in a simpler manner. In the majority of the cited tissue models explicitly based on turgor (like [Bassel et al., 2014](#), [Bessonov et al., 2013](#), [Boudon et al., 2015](#), [Bozorg et al., 2014, 2016](#), [Fozard et al., 2013](#), [Hamant et al., 2008](#), [Hervieux et al., 2016](#), [Hong et al., 2016](#), [Louveaux et al., 2016](#), [Robinson and Kuhlemeier, 2018](#), [Rudge and Haseloff, 2005](#), [Sampathkumar et al., 2014a](#)), steady and/or uniform cell turgor pressure was assumed, to simplify numerical analysis, and in absence of clear evidence of heterogeneities in cell pressures ([Long et al., 2018](#)). Typically, the cited authors have neglected the incompressibility of the liquid phase, which numerically would add a stiffening constraint that can be delicate to deal with. Very recently, authors have started reexamining the steady and uniform pressure hypothesis on both experimental ([Long et al., 2018](#)) and

theoretical (Cheddadi et al., 2019) grounds, using a 2D vertex-based model (and extending the discussion initiated by Lockhart, 1965, Ortega, 1985, to a multicellular context). This work is a novel step towards a multiphysical understanding of plant tissue mechanics, coupling both wall *viscoelastoplasticity* and water fluxes, and by doing so, allowing a more realistic treatment of pressure and incompressibility.

1.3.2 Computing growth

1.3.2.1 General implementation of growth

As soon as equilibrium is reached (see previous section), growth may be accounted for by modifying the rest configuration of the walls, *via* a modification of the growth tensor \mathbf{F}_g (Section 1.2.2). Due to Equations (1.6) and (1.7), this operation modifies the elastic deformation \mathbf{F}_e which generally results in a loss of equilibrium.

In one dimension, growth is defined as an increase of rest length as illustrated in Fig. 1.4(c). Such an approach has been used in several aforementioned vertex-based computational frameworks (*i.e.* Bessonov et al., 2013, Corson et al., 2009, Hamant et al., 2008, Rudge and Haseloff, 2005, Sapala et al., 2018) where growth is represented as an increase of the rest length of cell edges. Note that the implementation of growth in Sapala et al. (2018) does not completely comply with a morphoelastic description, as growth is not explicitly implemented as a modification of the rest lengths, but primarily as a displacement (rest length is redefined by the current lengths after displacement is applied, then elastic equilibrium due to pressure is computed). This geometric description is rather *ad hoc* and not based on physical considerations.

Other more abstract 2D and 3D Hamiltonian approaches have represented growth as an increase of cell (or tissue compartment) reference volume or shape (Alim et al., 2012, Cieslak et al., 2016, Merks et al., 2011).

As an alternative to vertex-based models, other authors have considered the growth of a multidimensional material, in general in 2D (or 3D in Kennaway et al., 2011). To do so, Bassel and coworkers have extended 1D edge growth to 2D finite elements (Bassel et al., 2014). Their solution consists of formulating growth as an increase of the rest length of the element edges, resulting in a redefinition of the element rest shape. Such a computational solution does not derive from an analytic continuum formulation and it precludes analytic treatments. In addition its physical and geometric interpretations are unobvious. This is because growth is here still seen as a 1D process, although the elastic regime is described through a proper multidimensional finite element formulation.

Bozorg and coworkers also pointed out the inadequacy of this approach in producing mesh-independent solutions and in properly representing mechanical anisotropies (Bozorg et al., 2016).

By contrast, the tensorial formulation of *morphoelasticity* (exposed in Section 1.2.2) provides a mathematically sounder, genuinely multidimensional extension of the 1D notion of rest length. This method has seen many applications to the growth of biological organisms (see Menzel and Kuhl, 2012, and references therein), and plant growth in particular, like in Boudon et al. (2015), Bozorg et al. (2016).

In this paragraph, I have discussed the procedures that are employed to define growth, from a strictly algorithmic point of view. In the sequel, I discuss the kinetic and physical assumptions that are made to govern the evolution of the rest configuration. These can be classified in two main views. In a first view, growth is assumed to be driven by *morphogens* (defined later). In another view, growth is seen as a mechanical relaxation process fueled by pressure-induced loading.

1.3.2.2 Growth as an explicit function of morphogens

How the rest configuration changes as time progresses depends upon the genetic regulatory network. To express this dependency, Coen and colleagues have proposed a parsimonious mechanism, suggesting that genes directly modulate the kinematic parameters of tissue growth (Coen et al., 2004, Green et al., 2010, Kennaway et al., 2011, Kuchen et al., 2012). This is done by expressing the growth tensor as a function of local gene expression and other abstract polarizing factors (that provide directional information):

$$\mathbf{L}_g = \mathbf{L}_g(\text{genes, polarizing factors}). \quad (1.9)$$

More generally, this approach consists in seeing growth as a *direct* function of *morphogens*⁷, that are abstractly defined as any substance that locally, and more or less directly, controls growth.

Similarly, the growth of the element edges in Bassel et al. (2014) was specified as a function of some morphogen concentration. Analogous solutions were used in Alim et al. (2012), Louveaux et al. (2016), Merks et al. (2011), Sapala et al. (2018) where growth was expressed as a function of abstract kinematic parameters, yet without explicitly invoking a morphogen.

The models cited in this section neither completely model the mechanical forces that cause tissue dilation, nor the rheological response of the tissue to these forces. The

⁷This term was coined by Alan Turing in “*The chemical basis of morphogenesis*” (Turing, 1952).

control of growth either relies on abstract biochemical parameters (Bassel et al., 2014, Kennaway et al., 2011) or phenomenological evolution laws (Alim et al., 2012, Louveaux et al., 2016, Merks et al., 2011, Sapala et al., 2018). It was yet established, that plant cell expansion is due to pressure-induced forces, that cause the wall to expand, with a speed that depends upon cell rheological properties (see Ali et al., 2014, Hamant and Traas, 2010, and references therein).

1.3.2.3 Turgor-driven growth

To mechanistically bridge the gap between genes and shape change, another approach was proposed, where genetic and metabolic activities regulate cell expansion *indirectly*, by modulating the biomechanical parameters of the cells (e.g. pressure, stiffness, etc.). In this approach, wall growth is explicitly seen as a response to forces, which are the product of turgor pressure.

In the approaches cited in Section 1.3.2.2, growth was decoupled from the elastic regime, and occurred independent of strain. This modality of growth can be physically seen as the result of some insertion of matter, that cause the tissue to dilate (which, in plants, does not occur independent of strain, Cosgrove, 2005). By contrast, turgor-induced growth is closer to a creep process, in which loading causes irreversible expansion provided a sufficient stretch is reached. At the scale of the cell wall, these two views are clearly in conflict. However, one cannot easily exclude the possibility that morphogen-driven growth at a continuous, supracellular scale may be a homogenization of pressure-driven growth at wall scale. For example, a morphogen in such an approach could be the local pressure itself.

More generally, these two views may in principle give rise to utterly different behaviors. In particular, their behavior with respect to residual stresses may differ, as proposed by Bozorg et al., who suggested that morphogen-driven growth theoretically favors large tensional and compressive residual stresses, compared to strain or stress-based laws (Bozorg et al., 2016). Figure 1.6 illustrates this principle considering a 1D beam undergoing either strain-independent (Fig. 1.6(a)) or strain-dependent (Fig. 1.6(b)) growth. In this example, strain-based growth favors a decrease of the elastic strain and a dissipation of strain energy. By contrast, strain-independent growth induces a compression, possibly resulting in a mechanical instability (*buckling*). Such compression cannot occur in the first case, as plastic expansion will not occur in absence of tension.

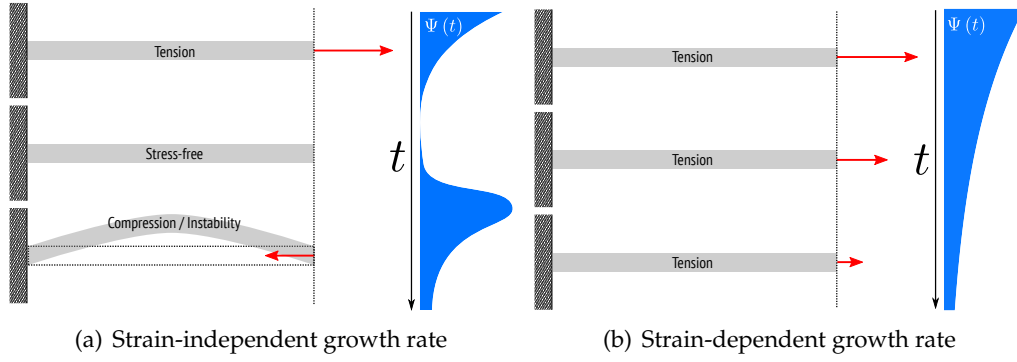


Figure 1.6: (a) Strain-independent and (b) strain-dependent growth of a beam. Red arrows indicate the force that is required to maintain the extremity of the beam at a constant position. Right plots indicate associated strain energy profiles (qualitative). In (a), the growth may favor the accumulation of compressive stress, that eventually may result in mechanical instabilities. The beam may buckle under compression, reaching a more energetically-favorable state. This does not occur in case (b), as growth stops in absence of strain.

The accumulation of residual stresses was dodged – rather artificially – in several strain-independent approaches. In [Kennaway et al. \(2011\)](#) the stresses ensuring tissue connectivity were canceled after the application of growth. In the 2D model detailed in [Sapala et al. \(2018\)](#), edge stretch was canceled after growth displacement was applied, leading to a new, stress-free configuration. This can be summarized by the following operation:

$$\forall n > 0 : \quad \mathbf{F}_g^n \leftarrow \mathbf{F}^{n-1} \quad (1.10)$$

that, by virtue of Eq. (1.6), is equivalent to a cancellation of the elastic deformation (n refers to the growth iteration).

In [Bonazzi et al. \(2014\)](#), [Hervieux et al. \(2016\)](#), [Hong et al. \(2016\)](#), growth was modeled through a similar stress cancellation, although the model used in these three works is explicitly strain-based. In fact, it is actually the simplest instance of a strain-based growth, which assumes fast and total relaxation of the strain at each growth step.

By construction, such a stress cancellation operation – which is justified neither theoretically nor experimentally – will fail in capturing the residual stresses that are experienced by the walls. In reality, as growth occurs nonhomogeneously and more or less fast, residual stresses may exist over long time periods, all the more that strain-based growth occurs only above a sufficient stretch level ([Cleland, 1959](#), [Lockhart, 1965](#), [Probine and Preston, 1962](#)). Below, the wall remains under elastic deformation. The plant tissue as a whole is therefore more plausibly in a homeostatic stressed configuration, as evidenced by cutting experiments ([Dumais and Steele, 2000](#)) or using an adhesion-deficient mutant ([Verger et al., 2018](#)) for example. Models not relying on

a cancellation of stress – typically morphoelastic approaches, in the case of continua – naturally provide more appropriate tools to model the possible effect of residual stresses on growth.

Several rheological models of strain-based growth have been developed in the form of plastic rheological laws. In particular, the fact that cell growth occurs only above a sufficient stretch level has been captured through various threshold-based formulations. This approach was pioneered by Lockhart in the 60s, who proposed a simple piece-wise linear model of one-dimensional cell growth (Lockhart, 1965). This was later on completed by Ortega, who extended Lockhart’s work to transient elastic regime (Ortega, 1985).

Lockhart’s model describes the uniform elongation of a single turgid, cylindrical cell, resulting from the quasi-static balance between constant turgor pressure and wall stretch. The formulation of wall creep is equivalent to that of a Bingham plastic, *i.e.* a plastic material yielding linearly above a threshold tension (see Fig. 1.7(a) and Bingham, 1917, Steffe, 1996):

$$L_g = \Phi (\sigma - \sigma_y)_+ . \quad (1.11)$$

Here, L_g depicts the 1D rate of growth (Eq. (1.8)); Φ depicts the cell *extensibility* (the “rate of irreversible flow”, in Lockhart’s terminology), that can be interpreted as the inverse of the characteristic time of growth; σ depicts the wall stress (that, at steady regime, is proportional to turgor pressure); σ_y is the yield stress, above which cell lengthening occurs; $(\cdot)_+ = \max(\cdot, 0)$ is the ramp function.

Equation (1.11) provides a nonlinear rheological law capturing the two-regime cell behavior previously observed *in vivo* (Cleland, 1959, Probine and Preston, 1962). Note that such a phenomenological abstraction disregards the molecular processes underlying growth. From a structural point of view, the wall material is an entanglement of various polysaccharides. In growing, meristematic or differentiating cells, the walls contain mainly cellulose, pectins and hemicelluloses that combined together form a more or less elastic solid medium (Ali et al., 2014, Cosgrove, 2005, Wolf et al., 2012). Several hundreds of proteins involved in wall modification have been identified. Many enzymes like pectin methyl-esterases or xyloglucan transglycosylase/hydrolases (Armezani et al., 2018, Cosgrove, 1999), or proteins like expansins (Cosgrove, 2000), may disrupt the wall constituents, causing the wall to yield irreversibly. In growing cells, this yield is simultaneously compensated by synthesis and secretion of new components into the wall, maintaining its mechanical and structural integrity (Cosgrove, 2005). To provide a mechanistic link between macroscopic rheology and microscopic wall biochemistry, several molecular-scale models have been proposed, mechanistically accounting for Lockhart-like behaviors (Ali and Traas, 2016, Dyson et al., 2012).

Essentially, Lockhart's model, which is one-dimensional, does not capture the various degrees of possible growth anisotropy. Mechanically, anisotropic growth may result from the fact i) that the forces applied to an infinitesimal wall parcel are anisotropically distributed and ii) that the mechanical response of the wall to these forces is itself anisotropic, due to anisotropies in the wall structural properties (Boudaoud, 2010, Hamant and Traas, 2010).

Several of the above-cited planar (or surface) models of growth have used viscous laws (Dupuy et al., 2006, Fozard et al., 2013), or Lockhart-like nonlinear laws (Bessonov et al., 2013, Cheddadi et al., 2019, Corson et al., 2009, Hamant et al., 2008, Rudge and Haseloff, 2005) to express the growth of the cell edges. In this kind of approach, growth anisotropy can in principle be achieved by i/ affecting different rheological parameters to each cell edge (as performed in Hamant et al., 2008, Rudge and Haseloff, 2005); or ii/ by applying an anisotropic distribution of forces to the nodes.

Two-dimensional continuum modeling approaches, that more properly capture the elastic regime, have extended Lockhart-like approaches to multidimensional expansion, using a tensorial formalism (Boudon et al., 2015, Bozorg et al., 2016, Fozard et al., 2013). Using a mathematical expression reminiscent of Lockhart's formulation (Eq. (1.11)), Boudon and coworkers have written the rate of growth L_g as a piece-wise linear function of the (reversible) *Green-Lagrangian strain* \mathbf{E} :

$$\mathbf{L}_g = \Phi(\mathbf{E} - \mathbf{E}_y)_+, \quad (1.12)$$

where $\mathbf{E} \stackrel{\text{def.}}{=} \frac{1}{2}(\mathbf{F}_e^T \cdot \mathbf{F}_e - \mathbf{I})$; $\mathbf{E}_y \stackrel{\text{def.}}{=} E_y \mathbf{I}$ is a positive spherical tensor defining the strain threshold in all directions; $(\cdot)_+$ depicts the tensor ramp function defined in Appendix 1.A and illustrated by Fig. 1.7(b) (Boudon et al., 2015). This approach was extended in Bozorg (2016), where L_g was a piece-wise linear function of strain, stress or morphogen concentration.

The growth law proposed in Boudon et al. (2015) (Eq. (1.12)) is formulated in terms of strain. Hence, it differs from Lockhart's original formulation, that was expressed in term of stress (Eq. (1.11)). This subtlety is not very meaningful in 1D models with linear constitutive relations, but becomes relevant considering anisotropic multidimensional materials like plant walls (see next Section 1.4), where strain and stress eigendirections do not coincide in general. The choice of a strain-based rather than stress-based growth law is motivated by a number of observations suggesting that the main orientation of the elastic strain correlates with that of growth, although, to my knowledge, this was never assessed in a quantitative manner. In organs like stems, for example, growth is almost exclusively longitudinal. Yet, due to the cylindrical shape of the stem, tangential stress is most probably mainly circumferential (from theoretical considerations, see

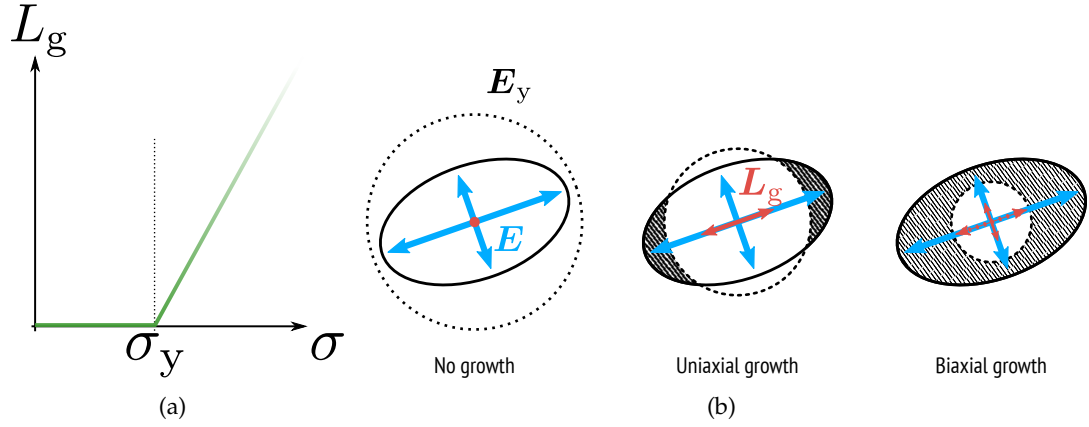


Figure 1.7: (a) Piece-wise linear regime corresponding to Eq. (1.11): growth rate vs stress. (b) Ellipse-based visualization of the growth multidimensional threshold in three cases. From left to right: all the eigenvalues of E are below the threshold (no growth occurs); one eigenvalue only exceeds the threshold (uniaxial growth); all eigenvalues exceeds the threshold (two-dimensional growth).

Landau and Lifshitz, 1986), while, anisotropic stiffness properties supposedly promote main strain in the longitudinal direction, *i.e.* orthogonal to main stress.

As Φ is a scalar, Eq. (1.12) states that the direction of main growth and elastic strain coincide, and that anisotropic growth mostly results from anisotropic elastic strain. Elastic strain in turn depends on stiffness properties and mechanical loading. To model this dependency, Boudon and colleagues have chosen a linear coupling between the *Green-Lagrangian strain* and the *second Piola-Kirchhoff stress* – that describes the internal forces in the reference configuration ($S = \frac{\partial \Psi}{\partial E}$) – through the fourth order *elasticity tensor* (\mathbb{C}_w , the subscript stands for 'wall'):

$$E = \mathbb{C}_w^{-1} : S \quad (1.13)$$

or equivalently $S = \mathbb{C}_w : E$ (where ':' depicts the tensor double-dot product). Tensor \mathbb{C}_w^{-1} is often referred as the *compliance tensor*.

We see next how Eqs. (1.12) and (1.13) mechanistically account for anisotropic growth, and how its parameters can be controlled.

1.4 Origin and regulation of anisotropic growth

In this section we ask where anisotropic growth originates from and what levers are accessible to a single cell to regulate it.

We may first remark that the right-hand side of Eq. (1.12) displays several parameters that may potentially be affected by the cell. Rheological parameters such as Φ and E_y

emerge from structural and physiological properties of the cell. These can be more or less directly controlled through genes. Yet, as they are scalars, they mostly affect the growth rate, and offer very little control upon growth anisotropy. From this statement, and by virtue of Eq. (1.13), we understand that growth anisotropy mostly depends upon the possible anisotropy of tensor quantities like \mathbb{C}_w and/or S .

It is generally assumed that stress largely results from tissue shape (Boudaoud, 2010), which affects the distribution of pressure forces. This idea is based on the common assumption that the plant tissue globally behaves like a pressurized shell (see Beauxamy et al., 2015, and references therein), indicating that local stress and geometry/curvature are coupled through a Young-Laplace type relationship. This assumption largely prevails in the literature, as it provides a tractable conceptual framework for inferring mechanical stress, and a convenient simplifying assumption for mathematical and computational modeling (e.g. in Bozorg et al., 2014, 2016, Hamant et al., 2008, Kierzkowski et al., 2012). The validity of this hypothesis however, was seldom assessed in a quantitative manner. Recently, computational modeling of multi-layered tissues led to suggest that the exact pattern of stress may in fact be not as simple as that expected from a simple shell-based consideration (Ali et al., 2019). In addition, the assumption that shape prescribes stress, no longer holds as soon as one considers the residual stresses arising from differential growth rates. Most plausibly, these stresses may be dominant in zones of highly heterogeneous growth rate, like the crease between the central zone of the SAM, and fast-growing incipient organs (Burian et al., 2013).

In principle, each individual cell is able to regulate its osmolarity, and by this means, to modify its internal pressure and the stress experienced by its walls. Yet, this kind of possible regulation offers limited control on the anisotropy of the cell wall stress, as it will mostly affect the amplitude of stress (that can be quantified by the trace of S). In addition, as mentioned before, an important part of cell stress is due to forces applied by the rest of the tissue (Boudon et al., 2015), which are transferred through cell adhesion (Verger et al., 2018). This leaves little room for a cell to autonomously regulate the anisotropy of its growth through stress.

By contrast, stiffness, which depends on the local structural and chemical properties of the wall, may be regulated by the cell in a more autonomous and direct manner. In particular, the main load-bearing element of the cell wall is its network of *cellulose microfibrils* (CMFs), which are stiff strand-like polymers (for reviews, see for example Ali et al., 2014, Hamant and Traas, 2010). These may be polymerized in disorganized directions, conferring isotropic stiffness to the wall (Fig. 1.9(a)); or around a preferential axis, resulting in more or less anisotropic stiffness (Fig. 1.9(b)). As captured by

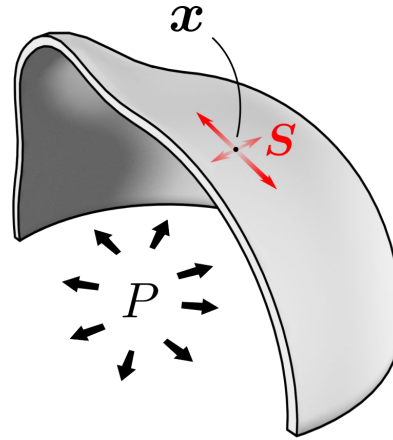


Figure 1.8: In the pressure vessel assumption, the tissue is equivalent to a pressurized membrane, on which one can define a tangent stress field. The situation is as if the whole tissue behaved like a gigantic pressurized cell (Kutschera, 1991). Reprinted from Oliveri et al. (2019).

Eqs. (1.12) and (1.13) and Fig. 1.9(c), this anisotropic stiffness may result in anisotropic elastic strain, and then anisotropic growth.

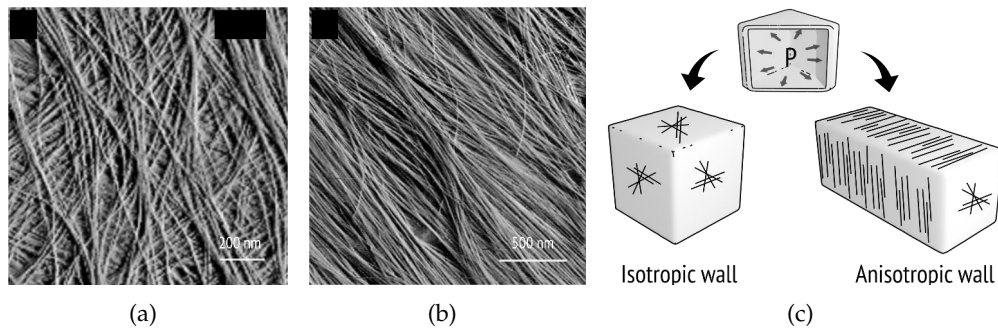


Figure 1.9: **Control of growth anisotropy through wall stiffness (a-b)** Atomic force microscopy scans of (a) disorganized and (b) organized cellulose microfibrils of onion cell wall. Adapted from Zhang et al. (2014) by permission from Springer. (c) The cell wall as a regulator of growth direction. Schematic of two pressurized cells with different wall properties and same initial shape. The first one has an isotropic wall and displays no privileged axis of growth. The other one displays hoop-shaped organization of microfibrils and expands axially. Reprinted from Oliveri et al. (2019).

CMFs are polymerized by transmembrane *cellulose synthase complexes* (CSCs) which move through the cell plasma membrane and synthesize cellulose in their wake (Kumar and Turner, 2015). In turn, this movement has been suggested to be guided by *cortical microtubules* (CMTs), as CSCs bind with CMTs (see Fig. 1.10(b) and Bringmann et al., 2012, Heath, 1974, Paredez et al., 2006). Microtubules are highly dynamic filamentous cytoplasmic polymers (Fig. 1.10(a)) that, collectively may come in disorganized arrangements, or in highly aligned bundles. This, and the functional association between

CMTs and CSCs, suggest that CMT organization prescribes CMF organization. What directional cue regulates the coalignment of CMTs?

At first thought, it seems logical to propose that this regulation depends on local gene expression, through the control of some chemical concentration. As a concentration is a scalar quantity, potentially only a chemical gradient could serve as a directional cue. Yet, to my knowledge, no scenario going in that direction was identified.

Alternatively, some intrinsic chemical polarity of cells would be a natural candidate for orienting CMTs. Several proteins – e.g. PIN1 (Barbier de Reuille et al., 2005, Jönsson et al., 2006, Smith et al., 2006) or BASL (Mansfield et al., 2018) – are asymmetrically distributed within a given cell, defining a polarity axis. At global scale, cells may also display cohesive polarity, making cell polarity a good candidate for an indicator of growth direction. Nonetheless, to date, no clear functional association was identified between these proteins and CMTs. In particular, while the alignment of CMT arrays may strongly correlate with PIN1 distribution, experimental results suggest the absence of a causal dependency between these, opening up the possibility of a third upstream factor (Heisler et al., 2010).

This factor could be mechanical stress (Heisler et al., 2010, Sampathkumar et al., 2014b), that in principle may provide a rich, multidimensional information field (three-component information: direction, anisotropy, amplitude). *In vivo*, CMT alignment strongly correlates with the theoretically predicted main axis of stress (Burian et al., 2013, Hamant et al., 2008, Hejnowicz et al., 2000, Robinson and Kuhlemeier, 2018, Sampathkumar et al., 2014a, Uyttewaal et al., 2012, Williamson, 1990). This was evidenced at different scales, on normal tissues – e.g. at the SAM epidermis (Burian et al., 2013, Hamant et al., 2008) or on pavement cell periclinal membranes (Sampathkumar et al., 2014a, Sapala et al., 2018) – as well as through more invasive experiments like cell ablation (Hamant et al., 2008, Sampathkumar et al., 2014a, Uyttewaal et al., 2012, Verger et al., 2018) or tissue compression (Hamant et al., 2008, Robinson and Kuhlemeier, 2018, Uyttewaal et al., 2012).

If mechanical stress feeds back on CMTs, then in turn, as CMTs affect CMF distribution in the wall, stress indirectly instructs the local growth pattern and controls morphogenesis (Hamant et al., 2008).

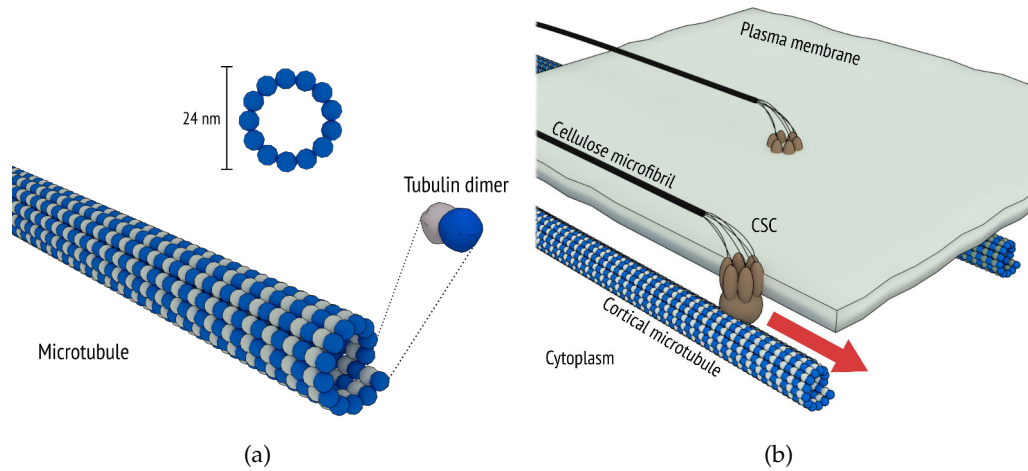


Figure 1.10: (a) Cartoon of the microtubule general structure. *Redrawn from T. Splettstößer, Wikimedia Commons.* (b) Cartoon showing the microtubule-guided transport of cellulose synthase complexes CSCs. The CSCs move along cortical microtubules and across the plasma membrane, and synthesize microfibrils in their wake. *Reprinted from Oliveri et al. (2019).*

1.5 Understanding the emergent behavior of stress-adaptive growing tissues

1.5.1 Literature review

From an abstract point of view, this kind of control typically characterizes a nonlinear dynamical system in the form of a feedback loop (illustrated in Fig. 1.11). This so-called stress feedback hypothesis consists of a local rule governing cell behavior. In turn, such local behavior integrates at larger scale into a global emergent behavior. To date, this emergent behavior, that is highly nonintuitive, is not fully characterized.

The stress feedback hypothesis is still under debate as the precise mechanism remains elusive, and because experimental proofs are often indirect (see Hamant et al., 2019). Modeling approaches, integrating the stress feedback as a working hypothesis, may allow assessing its theoretical plausibility and implications on morphogenesis, notably by yielding testable predictions.

Several growth models have integrated a stress-based control of the anisotropy of stiffness. This was done through 2D spring-based modeling (Hamant et al., 2008, Sapala et al., 2018), in 2D generalized Hamiltonian approaches (Alim et al., 2012, Uyttewaal et al., 2012), as well as in continuum approaches (Bozorg, 2016, Hervieux et al., 2016). Other authors have used a static model (without growth) to model the stress feedback in the elastic regime (Bozorg et al., 2014).

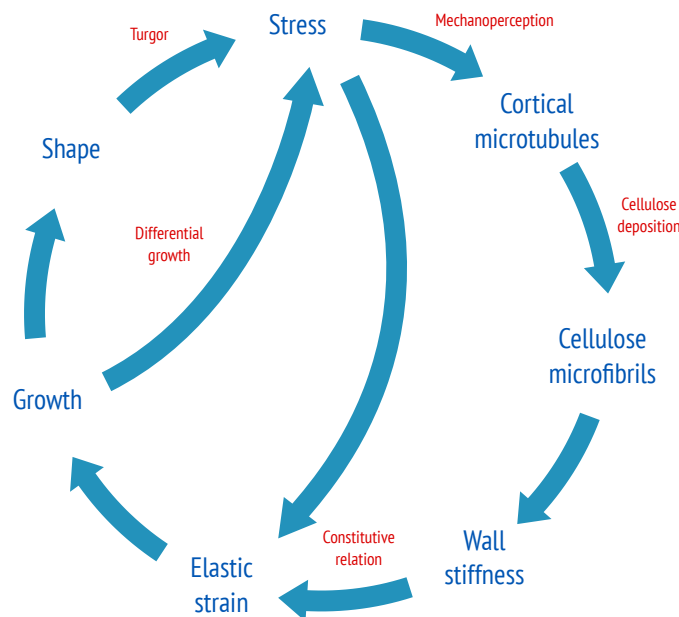


Figure 1.11: Hypothetical feedback loop governing plant morphogenesis: stress feeds back on CMTs, and regulates growth.

In [Sapala et al. \(2018\)](#), CMTs were modeled in a discrete manner, by representing each CMT as a spring connecting two nodes of a given 2D cell polygon. Essentially, at each iteration, each node is considered as the potential starting point of a CMT, that is inserted only if a number of geometric conditions are verified. This method is relatively simple but rather *ad hoc*; in particular, the number of CMTs (therefore global stiffness) may depend on mesh resolution. In [Hamant et al. \(2008\)](#), where cells were modeled as a polygon made of springs, the spring constants were modulated by projecting the main stress axis (obtained as the circular mean of the node forces, see [Mardia, 1972](#)) on each edge. In the 2D Hamiltonian approach detailed in [Alim et al. \(2012\)](#), the rate of growth defined for each cell is a function of the cell stress tensor, hence growth does not explicitly result from a change in the cell rheological properties. Finally, in other continuum approaches such as [Bozorg et al. \(2014\)](#), [Hervieux et al. \(2016\)](#) stiffness coefficients directly depend upon various observables of the stress tensor (*i.e.* trace, anisotropy, main orientation).

In [Alim et al. \(2012\)](#), [Bozorg et al. \(2014\)](#), [Hamant et al. \(2008\)](#), [Hervieux et al. \(2016\)](#), the link between stiffness and stress is modeled in a more or less phenomenological manner. These works in particular do not consider the dynamics and organization of underlying subcellular actors such as CMTs or CMFs. Therefore, the parameters (stiffness, sensitivity of the stress feedback) of these models are abstract, and not connected to the chemical constants that govern the reorganization of CMTs and CMFs. To advance towards a more quantitative understanding of plant growth, finer, multiscale approaches are required.

Agent-based models have also been developed, where microtubules are represented as individual agents (Allard et al., 2010, Chakraborty et al., 2018, Dixit and Cyr, 2004, Mirabet et al., 2018, Muratov and Baulin, 2015, Nédélec, 2002, Tindemans et al., 2010). This kind of approach allows to finely study the dynamics of self-organization of microtubules, that rely mostly on mutual short-range microtubule-microtubule interactions. However, as they focus on complex behaviors at molecular scale, involving an important number of microtubules, extending these models to virtual tissues composed of several hundreds of cells would be impracticable.

1.5.2 Objectives of the thesis

Continuum approaches provide a mathematically tractable framework that benefits from a huge arsenal of analytic and numerical techniques. However integrating small scale molecular features within such framework can be challenging. In this thesis I couple continuum mechanics with small scale molecular dynamics, proposing a parsimonious, coarse-grained model of the wall structure and stress-based CMT organization.

To some extent, this thesis was influenced by the reflections developed in Bozorg (2016), where a pioneering approach of the problem is proposed, interrogating the macroscopic properties of the stress feedback and its potential role in the large context of morphogenesis. Along a similar line, viewing biological development as a nonlinear dynamical system, I here ask a number of general questions. What are the emergent properties of a growing material whose growth is itself controlled through the mechanical stress field? What stable shapes are allowed by this system? Under what conditions? Would growth be globally over-constrained if all walls stiffened along the main stress?

To answer these questions, I first developed a parsimonious, multiscale theoretical model of a stress-responsive tissue (Chapter 2). This model establishes a dependency between the chemical kinetics of CMTs/CMFs, and continuum tissue mechanics, in a more explicit manner than in previous approaches. Analytic and numerical tissue-scale FEM studies were performed, in static regime (without growth). In this model, stiffness is a function of the CMF distribution, that itself results from CMT-guided cellulose deposition. CMT distribution in turn depends upon the stress that modulates their chemical stability, in a phenomenological manner (since no mechanism has been actually identified).

In Chapter 3, the model is coupled with growth, previously implemented in [Boudon et al. \(2015\)](#). This model provides a computational framework in which growth behavior is connected to physical quantities and chemical constants. In particular, growth affects the density and orientation of the fibers (that are transported by the growing medium), and therefore feeds back on stiffness. This growth model is then applied to a practical biological problem in Chapter 4, suggesting that stress feedback may be a mechanism for the amplification and maintenance of incipient leaf flatness.

Appendices

1.A The tensor ramp function

Here, the scalar ramp function $(\cdot)_+ \stackrel{\text{def.}}{=} \max(\cdot, 0)$ is extended to the set of the symmetric second order tensors of dimension d . Let \mathbf{T} be such a tensor; \mathbf{T} admits a unique rank decomposition, that reads:

$$\mathbf{T} = \sum_{n=1}^d \lambda_n \mathbf{t}_n \otimes \mathbf{t}_n \quad (1.14)$$

where $\{\lambda_n\}_{n \in [1,d]} \subset \mathbb{R}$ depicts the set of the eigenvalues of \mathbf{T} , and $\{\mathbf{t}_n\}_{n \in [1,d]} \subset \mathbb{R}^d$ is an orthonormal family of corresponding eigenvectors. We define the tensor ramp function as:

$$(\mathbf{T})_+ \stackrel{\text{def.}}{=} \sum_{n=1}^d (\lambda_n)_+ \mathbf{t}_n \otimes \mathbf{t}_n. \quad (1.15)$$

This definition assures that $(\mathbf{T})_+$ is a positive symmetric tensor, since all its eigenvalues are positive.

Chapter 2

Regulation of plant cell wall stiffness by mechanical stress: a mesoscale physical model

Foreword

This is a post-peer-review, pre-copyedit version of an article published in *Journal of Mathematical Biology* (here referred as [Oliveri et al., 2019](#)). The final authenticated version is available online at doi.org/10.1007/s00285-018-1286-y. Several minor corrections have been added to this chapter with respect to the published version.

As the first author of this publication, I am the main contributor to the work hereafter exposed, and wrote most part of the manuscript.

2.1 Abstract

A crucial question in developmental biology is how cell growth is coordinated in living tissue to generate complex and reproducible shapes. We address this issue here in plants, where stiff extracellular walls prevent cell migration and morphogenesis mostly results from growth driven by turgor pressure. How cells grow in response to pressure partly depends on the mechanical properties of their walls, which are generally heterogeneous, anisotropic and dynamic. The active control of these properties is therefore a cornerstone of plant morphogenesis.

Here, we focus on wall stiffness, which is under the control of both molecular and mechanical signaling. Indeed, in plant tissues, the balance between turgor and cell wall elasticity generates a tissue-wide stress field. Within cells, mechano-sensitive structures, such as cortical microtubules, adapt their behavior accordingly and locally influence cell wall remodeling dynamics. To fully apprehend the properties of this feedback loop, modeling approaches are indispensable. To that end, several modeling tools in the form of virtual tissues have been developed. However, these models often relate mechanical stress and cell wall stiffness in relatively abstract manners, where the molecular specificities of the various actors are not fully captured.

In this paper, we propose to refine this approach by including parsimonious biochemical and biomechanical properties of the main molecular actors involved. Through a coarse-grained approach and through finite element simulations, we study the role of stress-sensing microtubules on organ-scale mechanics.

2.2 Main quantities

L_g	Growth tensor
E	Green-Lagrangian elastic strain tensor
S	PK2 stress tensor
Φ	Cell wall extensibility
τ	Cell wall yield strain
\mathbb{C}_w	Cell wall stiffness tensor
\mathbb{C}_g	Stiffness tensor associated with the wall's isotropic matrix
\mathbb{C}_f	Stiffness tensor associated with CMFs
Y, ν	Wall matrix reduced Young's modulus and Poisson's ratio
Y_f	Microfibril reduced Young's modulus
θ	Angle parameter in the wall tangential plane
e_θ	Unit vector oriented by θ
Θ	Projector on span(e_θ)
$\rho(\theta)$	Angular density of CMFs
$\phi(\theta)$	Angular density of CMTs
$f(\theta)$	Angular density of force (per unit surface)
$\hat{\rho}_n, \rho_n, \tilde{\rho}_n$	Complex, even and odd Fourier coefficients of ρ
$\hat{\phi}_n$	Complex Fourier coefficients of ϕ
\hat{f}_n	Complex Fourier coefficients of f
α_ρ	Anisotropy of CMFs
α_ϕ	Anisotropy of CMTs
α_f	Anisotropy of forces
k_ρ, k'_ρ	Microfibril polymerization and depolymerization constants
k_ϕ	Microtubule polymerization constant
k'_ϕ	Inverse of stress-free microtubule half-life
γ	Coupling coefficient of the stress-induced microtubule stabilization
$k'_\phi(\theta) = k'_\phi e^{-\gamma f(\theta)}$	Angular microtubule depolymerization coefficient
c_0	Total concentration of tubulin
K_ρ, K_ϕ	Equilibrium constants of the CMF/CMT kinetics
η	Measure of the relative stiffness between the gel and the fiber

2.3 Introduction

In the absence of cell migration, plant development occurs through inhomogeneous and anisotropic irreversible deformations of the tissue, driven by turgor-induced forces. It has been proposed that global shape changes are controlled locally, at the cellular

level, through three geometric parameters: growth rate, growth direction, and the degree of growth anisotropy (Coen et al., 2004, Erickson, 1976, Kennaway et al., 2011). How individual cells regulate these parameters is fundamental to understanding how plants control their shape.

Growth results from turgor-induced forces, that are counterbalanced by the cell wall, a stiff exoskeleton surrounding the cells and preventing them from bursting (Hamant and Traas, 2010). The wall is a complex assembly of cellulose microfibrils cross-linked by a network of polysaccharides, including hemicelluloses and pectins (Cosgrove, 2001). When stretched above a certain threshold these cross-links may break, allowing microfibrils to slide (as modeled in Dyson et al., 2012) thus causing the irreversible expansion of the wall. New material is simultaneously added to maintain wall integrity (Cosgrove, 2005). This irreversible process defines cell growth. Throughout development, cells may have different growth rates at different positions and times. In addition, since microfibrils can be deposited in more or less parallel arrays, and/or because the wall may be mechanically stressed unequally in various directions, cell growth may be anisotropic (Fig. 2.1(a), Baskin, 2005).

The central role of wall mechanics in growth has motivated the development of a *rheological* formalization of tissue expansion, initiated by Lockhart on one-dimensional (1D) cell expansion (Lockhart, 1965, Ortega, 1985). In brief, Lockhart modeled the cell wall as a Bingham plastic undergoing linear irreversible expansion above a certain yield pressure. In recent years, computational modeling has been used to represent higher-dimensional tissue expansion, in 2D or 3D (for a review, see Ali et al., 2014). Several studies were based on discrete tissue descriptions, where the tissue was modeled as a network of 1D elements (Corson et al., 2009, Dupuy et al., 2006, Hamant et al., 2008). This allows a reasonable trade-off between modeling expressiveness and computational complexity, but fails at representing efficiently a number of specific properties of higher-dimensional mechanics (Gelder, 1998), such as shear, incompressibility, and anisotropy. To alleviate this difficulty, other authors have adapted the formalism of continuous media to growth (Boudon et al., 2015, Bozorg et al., 2016, Dumais et al., 2006, Dyson and Jensen, 2010, Fozard et al., 2013, Goriely and Ben Amar, 2007, Hervieux et al., 2016, Rojas et al., 2011, Sassi et al., 2014). Boudon et al. and Bozorg et al. developed a Lockhart-like 2D description of growth using a second-order growth tensor (\mathbf{L}_g), expressed as a function of the elastic (reversible) deformation, represented by the elastic strain tensor (\mathbf{E}):

$$\mathbf{L}_g = \Phi(\mathbf{E} - \tau \mathbf{I})_+ \quad (2.1)$$

where Φ is the wall extensibility, τ is a threshold strain, $(\cdot)_+$ is the tensor ramp function (defined in Appendix 1.A) that allows growth only in directions where deformation is higher than the threshold τ ; and \mathbf{I} is the second-order identity tensor (Boudon et al., 2015, Bozorg et al., 2016). For the sake of simplicity, it is generally assumed that the cell wall extensibility (Φ) is quantified by a scalar coefficient or, equivalently, that growth and strain directions coincide. In the linear elasticity assumption (Hooke's law), strain and stress (depicted by tensor \mathbf{S}) are linearly coupled via the fourth-order elasticity tensor (\mathbb{C}_w):

$$\mathbf{E} = \mathbb{C}_w^{-1} : \mathbf{S} \quad (2.2)$$

(':' is the tensor double-dot product). Combining the two previous expressions shows that anisotropic growth results from an anisotropy in stress and/or in stiffness. There is a reduced potential for cells embedded in a tissue, to autonomously regulate their local stress, which is mostly imposed by the rest of the tissue (Boudon et al., 2015). By contrast, by modifying locally the wall's structure, cells can affect growth anisotropy in a more direct manner.

Such a local regulation does occur in plant tissues and has been proposed to rely partly on the activity of cortical microtubules. Microtubules participate in locally structuring the cell wall, by guiding the trajectories of the transmembrane complexes synthesizing the microfibrils (Fig. 2.1(b), Paredez et al., 2006). There are important indications, that microtubules in many cells align along the main stress direction (Hamant et al., 2008, Sampathkumar et al., 2014a,b, Williamson, 1990). This has led to the hypothesis that forces, generated in the tissues by turgor and differential growth rates, feed back into the structure of the cell wall via microtubule organization.

This so-called *stress feedback* has been integrated into tissue-scale models. Hamant et al. used a spring-based cell discretization of the tissue (Hamant et al., 2008). In this approach, stress feedback is taken into account by modulating spring constants according to the angle between the spring and the cell's average loading axis. Other authors have instead used a continuous formalism, expressing a coupling between multidimensional stiffness and stress, in Bozorg et al. (2014), through triangular biquadratic springs (Delingette, 2008) and in Hervieux et al. (2016), through the finite element method (FEM). In the last two implementations, stress is considered as an input-parameter that modulates the main axis of stiffness and its anisotropy.

These studies express an abstract phenomenological link between stress and stiffness, not explicitly taking into account the underlying molecular mechanisms. Here, we describe a modeling approach where subcellular processes involved in the wall synthesis are more explicitly described. Based on a parsimonious kinetic model of microtubule reorganization and microfibril synthesis, we propose a homogenized expression of the

wall's dynamical elastic properties. We present a general expression of force-induced microtubule stabilization in the direction of higher tension. This simple model leads to subtle predictions on the dynamics of microtubule arrays and gives a quantitative description of wall stiffness. We derive an analytical, quantitative behavior of the microtubule cortex and the material mechanics. Tissue-scale FEM simulations of the system reproduced several macroscale behaviors previously observed *in vivo*.

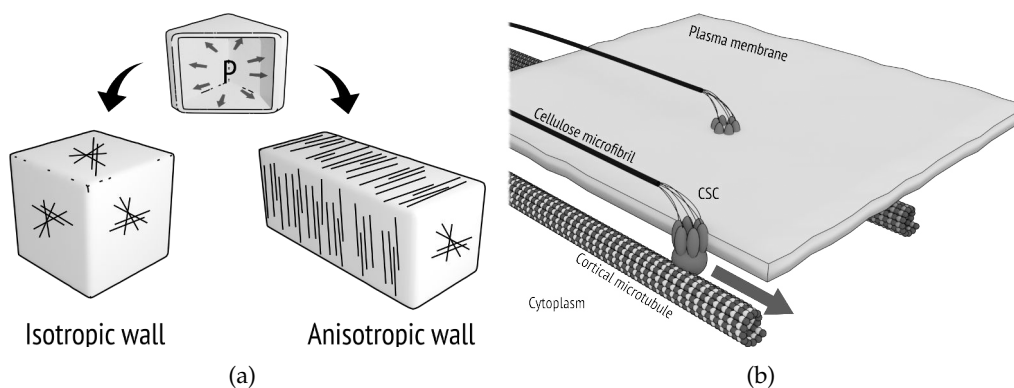


Figure 2.1: (a) The cell wall as a regulator of growth direction. Schematic of two pressurized cells with different wall properties and same initial shape. The first one has an isotropic wall and display no privileged axis of growth. The other one displays hoop-shaped organization of CMFs and expands axially. (b) Cartoon showing the CMT-guided transport of cellulose synthase complexes (CSCs). CSCs move along cortical microtubules and across the plasma membrane, and synthesize microfibrils in their wake. Redrawn after Landrein and Hamant (2013a).

2.4 Model

In this section, we introduce our model theoretically, by first detailing the elastic behavior of the wall considered as a fiber-reinforced medium, and then the kinetics of CMTs and CMFs.

2.4.1 Mechanical model

2.4.1.1 Mechanical equilibrium

The meristem is generally assimilated to a thin pressurized shell with uniform thickness (Fig. 2.2(a), Beuzamy et al., 2015, Kutschera, 1991). This shell defines a continuum on which stiffness, strain and stress can be defined as tensor fields evolving in space and time, respectively $\mathbb{C}_w(\mathbf{x}, t)$, $\mathbf{E}(\mathbf{x}, t)$ and $\mathbf{S}(\mathbf{x}, t)$ (NB, since the next developments are made at a fixed position \mathbf{x} , we get rid of the spatial parameter \mathbf{x} in the following

notations). Due to large radius of curvature (compared to thickness), we assume plane stress within the surface (Landau and Lifshitz, 1986).

Because biological processes occur at a time scale much smaller than elastic relaxation, we assume quasi-static mechanical equilibrium at any time:

$$\forall t \quad \nabla \cdot \boldsymbol{\sigma}(t) + \mathbf{f}_{\text{ext}}(t) = \mathbf{0} \quad (2.3)$$

where $\nabla \cdot \boldsymbol{\sigma}$ is the divergence of the tangential *Cauchy stress* and \mathbf{f}_{ext} is the density of pressure force applied normally to the surface at \mathbf{x} .

2.4.1.2 Constitutive relation of the material

In higher plants, the wall is largely composed of three polysaccharides: pectins, hemicelluloses and cellulose (Cosgrove, 2001). The former is polymerized as stiff, linear strands, the CMFs. They are embedded in a matrix of pectins and hemicelluloses that form an isotropic, soft hydrogel. In addition, other polysaccharides, e.g. hemicellulose, as well as numerous enzymes constantly interact and remodel these two main structural elements (Wolf et al., 2012).

In the sequel, it is assumed that this fiber-reinforced hydrogel behaves as a linear deformable solid (Eq. (2.2)), composed of an isotropic homogeneous matrix and a fiber network (Fig. 2.2(b)). We assume that fibers are much stiffer than the rest of the material. For that, we neglect the mechanical effect of the possible interaction between the fibers and the matrix. The whole system (*i.e.* fibers + matrix) can thereby be seen as the parallel association of both components. The resulting elasticity tensor \mathbb{C}_w is therefore the sum of the fiber-related \mathbb{C}_f and the matrix \mathbb{C}_g :

$$\mathbb{C}_w = \mathbb{C}_g + \mathbb{C}_f. \quad (2.4)$$

The gel is assimilated to an isotropic deformable solid, described by a Young's modulus and a Poisson's ratio. Tensor \mathbb{C}_g can be expressed in an arbitrary surface basis ($\mathbf{e}_x, \mathbf{e}_y$), by employing the Voigt form¹

$$[\mathbb{C}_g] = Y \begin{bmatrix} 1 & \nu & 0 \\ \nu & 1 & 0 \\ 0 & 0 & \frac{1-\nu}{2} \end{bmatrix} \quad (2.5)$$

¹Due to the symmetries $\mathbb{C}_{ijkl} = \mathbb{C}_{klij}$ and $\mathbb{C}_{ijkl} = \mathbb{C}_{ijlk} = \mathbb{C}_{jikl} = \mathbb{C}_{jilk}$, a given elasticity tensor \mathbb{C} can be expressed as a 3×3 matrix (Voigt notation).

where ν and Y respectively stand for the Poisson's ratio and for the reduced Young's modulus².

Describing the fiber-related component, which contains the anisotropy of the whole material, requires a mechanical model of the fibers. Those are modeled by linear springs distributed in the surface's tangential plane. For the sake of simplicity, we assume that they do not mutually interact and do not move relative to the medium³.

Let $\rho(\theta)$ be the local angular density of fibers⁴ in an arbitrary tangential reference frame (e_x, e_y) (Fig. 2.2(b)). Since mechanically, a fiber is apolar⁵, function ρ is π -periodic. As one may expect, the elasticity tensor \mathbb{C}_f is a function of the density ρ . By assuming affine deformation of the fibers (namely that each individual fiber is deformed as the macroscopic medium), one shows that \mathbb{C}_f can be expressed according to the low-frequency Fourier coefficients $\rho_0, \rho_1, \tilde{\rho}_1, \rho_2$ and $\tilde{\rho}_2$ (Appendix 2.A and [Cox, 1952](#)):

$$[\mathbb{C}_f] = \frac{\pi Y_f \rho_0}{16} \begin{bmatrix} 3 + \frac{\rho_2 + 4\rho_1}{\rho_0} & 1 - \frac{\rho_2}{\rho_0} & \frac{2\tilde{\rho}_1 + \tilde{\rho}_2}{\rho_0} \\ 1 - \frac{\rho_2}{\rho_0} & 3 + \frac{\rho_2 - 4\rho_1}{\rho_0} & \frac{2\tilde{\rho}_1 - \tilde{\rho}_2}{\rho_0} \\ \frac{2\tilde{\rho}_1 + \tilde{\rho}_2}{\rho_0} & \frac{2\tilde{\rho}_1 - \tilde{\rho}_2}{\rho_0} & 1 - \frac{\rho_2}{\rho_0} \end{bmatrix} \quad (2.6)$$

where Y_f is the reduced Young's modulus of one fiber.

Assuming that the wall's thickness, the gel's stiffness and Y_f are constant in time, Eqs. (2.4) to (2.6) imply that the evolution of the wall's mechanical properties only results from the dynamics of the CMFs distribution.

2.4.2 The dynamics of cortical microtubules and microfibrils

The orientation of cellulose deposition depends on the orientation of cortical microtubules that guide the trajectories of transmembrane cellulose synthase complexes (CSCs). In our model, tubulin (namely, the elementary molecular sub-unit of CMTs) is described as a biphasic gel: it can be found either in a monomeric soluble state or in a polymerized form (the CMT). The distribution of CMTs is described through its π -periodic angular distribution ϕ .

In our model, CMT orientation influences only the orientation of cellulose deposition, and has no impact on the mass of cellulose polymerized per unit time. In brief, this

²The actual Young's modulus divided by $1 - \nu^2$.

³In reality, fibers are connected through the hemicellulose network, that is yet assumed to be significantly softer than the fibers and then negligible.

⁴For given positions x and angle θ (angle with e_x in the tangential plane), $\rho(\theta) d\theta$ is the number of fibers at the vicinity of x whose orientation angle lies in the infinitesimal interval $[\theta, \theta + d\theta]$.

⁵A fiber can be modeled by a straight line, equivalently defined by the unit vectors $e_\theta = \cos \theta e_x + \sin \theta e_y$ or $-e_\theta$.

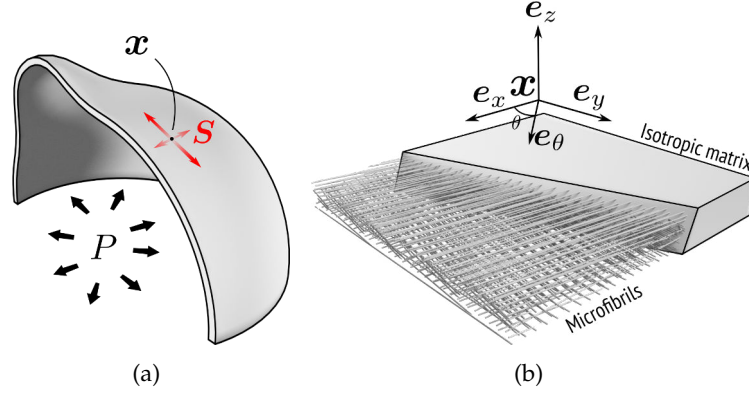


Figure 2.2: **Mechanical model of the epidermis.** (a) The tissue is represented by a pressure-filled thin membrane. (b) Sketch of the wall structural model: CMFs are immersed in an elastic solid isotropic hydrogel.

relies on the hypothesis that a CSC protein participates to cellulose deposition regardless of whether it is bound to a CMT (Emons et al., 2007), as detailed in Appendix 2.B. Hence, cellulose polymerization is expressed as a function of the *normalized* microtubule distribution $\phi^* = \phi / \int_{\pi} \phi$. In addition to that polymerization term, we introduce an isotropic decay that prevents unlimited accumulation of cellulose in the wall. At the considered time scale, we neglect growth, and then, the CMF advection that it would induce. Finally, we express the evolution of the angular density of CMFs as:

$$d_t \rho(\theta, t) = k_{\rho} \phi^*(\theta, t) - k'_{\rho} \rho(\theta, t), \quad (2.7)$$

(k_{ρ} and k'_{ρ} are kinetic constants).

The evolution of the microtubule angular density ϕ emerges from a description of the tubulin polymerization/depolymerization kinetics. Tubulin is present in the cytoplasm under two forms: a soluble one composed of free dimers (with concentration c_{free}), and a polymerized one depicted by ϕ . We assume that its total concentration c_0 (namely regardless of its form) is constant and uniform. This concentration verifies:

$$c_0 = c_{\text{free}} + \int_{\pi} \phi. \quad (2.8)$$

Microtubule dynamics is governed by the conservation equation of tubulin:

$$d_t \phi(\theta, t) = k_{\phi} c_{\text{free}} - k'_{\phi}(\mathbf{S}, \theta) \phi(\theta, t). \quad (2.9)$$

The synthesis term (first term on the right-hand side) describes the isotropic CMT nucleation from the pool of free dimers (with kinetic constant k_ϕ). The second term models the microtubule depolymerization rate. Unlike in Eq. (2.7), the microtubule depolymerization rate varies with the orientation according to the angular function $k'_\phi(\mathbf{S}, \theta)$, that models stress sensing. The last is described by assuming that microtubule stability increases with the tensile stress angular distribution f , reducing the degradation probability according to:

$$k'_\phi(\mathbf{S}, \theta) = k'_\phi e^{-\gamma f(\mathbf{S}, \theta)}, \quad (2.10)$$

where k'_ϕ stands for the inverse half-life of polymerized tubulin in absence of stress, γ is a (positive) coupling parameter, and $f(\mathbf{S}, \theta) = \mathbf{e}_\theta \cdot \mathbf{S} \cdot \mathbf{e}_\theta$ depicts the force per unit surface in direction θ (details on the establishment of Eq. (2.10) are provided in Appendix 2.C). This angular variability of the microtubule depolymerization probability over the local orientation of the stress field is the key feature of our modeling approach and accounts for the experimentally-observed tendency of microtubules to align in the direction of maximal tension (Hamant et al., 2008). The general hypothesis behind this is that microtubules under tension undergo some conformational change making them less sensitive to katanin severing activity and therefore increasing their lifetime (Landrein and Hamant, 2013a, Sampathkumar et al., 2014a).

In the last two sections, we established a dynamical model of the wall's stiffness, by describing (i) the relation between the elasticity tensor \mathbb{C}_w and the microfibril bundle (Eqs. (2.4) to (2.6)), (ii) the microtubule-guided deposition of CMFs (Eq. (2.7)) and (iii) the stress-regulated kinetics of CMTs (Eqs. (2.8) to (2.10)). Importantly, Eqs. (2.4) to (2.7) and (2.11) can be conceptually combined into a single ordinary differential equation relating the time evolution of the stiffness elasticity \mathbb{C}_w to stress.

2.4.3 Tissue-scale numerical simulation procedure

Our model expresses the *local* behavior of the system, focusing on an infinitesimal portion of tissue in which stress is an exogenous parameter. We extended the analysis to a large-scale tissue, represented as an elastic surface, loaded with constant pressure (Fig. 2.2(a)). In this context, stress results from curvature and pressure; and may evolve in time as the stiffness properties vary.

Due to its nonuniform curvature, the tissue experiences nonuniform stress field. In practice, the stress and the elastic response of such a system (its displacement field) can generally only be determined by numerical approximation.

For that, we employed the FEM (in space) with triangle meshes, coupled with a *finite difference* time discretization. At initial time $t = t_0$, the CMT/CMF system is considered at equilibrium (null-stress steady state, isotropic distribution, see Eq. (2.43)). The structures are not loaded and no residual stress is assumed.

Assuming that the CMTs' transient regime is much shorter than that of cellulose deposition, CMT distribution can be considered at equilibrium, which is described by the steady-state solution of Eq. (2.9):

$$\phi^\infty(\mathbf{S}, \theta) = \frac{c_0 K_\phi e^{\gamma f(\mathbf{S}, \theta)}}{1 + K_\phi \int_\pi d\theta' e^{\gamma f(\mathbf{S}, \theta')}}, \quad (2.11)$$

with $K_\phi = k_\phi/k'_\phi$, the equilibrium constant of the CMT stress-free polymerization process.

Under this assumption, only the CMF distribution can vary. It is iteratively updated, on each finite element, by considering the steady-state CMT distribution (Eq. (2.11)), and the CMF evolution equation (Eq. (2.7)) solved through Euler's method (Eq. (2.42)). After this step, stiffness is updated according to Eq. (2.6), bringing the system out of equilibrium and providing an new initial state. For further details on the numerical pipeline, refer to Appendix 2.E.

2.5 Results

We detail hereafter the outputs of the model exposed in the previous section. We first expose how the CMT steady state accounts for the experienced stress field. We then show how the cell wall equivalent elasticity tensor is modified and how that leads to nontrivial strain response.

2.5.1 The steady CMT/CMF pattern reflects stress

This section exposes results on the steady-regime behavior derived from Eqs. (2.4) to (2.7) and (2.11) (refer to Appendix 2.D for detailed mathematical developments).

The steady-state expression of the CMT distribution ϕ^∞ (Eq. (2.11)) expresses their tendency to crowd in the direction of largest tensile stress (Fig. 2.3(a)). Quantification of the microtubule coalignment has been proposed by Boudaoud et al. (2014), based on the concept of *nematic tensor* (see Section 2.E.3). The authors define *microtubule anisotropy* as the ratio:

$$\alpha_\phi^\infty = \frac{Q_{\phi,1}^\infty - Q_{\phi,2}^\infty}{Q_{\phi,1}^\infty + Q_{\phi,2}^\infty} \in [0, 1] \quad (2.12)$$

where $Q_{\phi,1}^\infty \geq Q_{\phi,2}^\infty \geq 0$ are the eigenvalues of the CMT nematic tensor. This definition of anisotropy can be expressed in the Fourier formalism, as it is equivalent to:

$$\alpha_\phi^\infty = \frac{|\hat{\phi}_1^\infty|}{\hat{\phi}_0^\infty} \left(= \frac{\sqrt{\phi_1^2 + \tilde{\phi}_1^2}}{\phi_0} \right) \quad (2.13)$$

where $\{\hat{\phi}_n^\infty\}_{n \in \mathbb{N}}$ depicts the Fourier sequence of ϕ^∞ (see Section 2.E.3 for a proof). Similarly, the anisotropy of stress can be defined as $\alpha_f = (S_1 - S_2) / (S_1 + S_2)$ (where $S_1 \geq S_2$ are the stress eigenvalues)⁶. Again, by developing the expression $f(\mathbf{S}, \theta) = \mathbf{e}_\theta \cdot \mathbf{S} \cdot \mathbf{e}_\theta$, one shows that $\alpha_f = 2|\hat{f}_1|/\hat{f}_0 (= 2\sqrt{f_1^2 + \tilde{f}_1^2}/f_0)$.

As shown in Figs. 2.3(a) and 2.3(b) and Eq. (2.35), α_ϕ^∞ is a growing function of the stress anisotropy, but also of the mean stress $f_0 = \text{Tr } \mathbf{S}/2$. Similarly, the concentration of CMTs ($\pi\phi_0^\infty/2$) is a growing function of stress anisotropy and amplitude (Fig. 2.3(c) and Eq. (2.33)). Note that if high stress amplitude favors high CMT coalignment, we can expect that this effect also comes with a lengthening of the transient regime's duration, due to the augmented stability of CMTs that diminishes CMT turnover⁷.

Considering infinitesimal displacements, one can assume that the global shape, and therefore the distribution of the pressure forces, do not vary during the evolution of the material stiffness. This allows to derive the steady-state CMF distribution from Eq. (2.11):

$$\rho^\infty(\mathbf{S}, \theta) = K_\rho \phi^{*\infty}(\mathbf{S}, \theta) = \frac{K_\rho e^{\gamma f(\mathbf{S}, \theta)}}{\int_\pi d\theta' e^{\gamma f(\mathbf{S}, \theta')}}. \quad (2.14)$$

The anisotropy of the CMF steady distribution, defined as $\alpha_\rho^\infty = |\hat{\rho}_1^\infty|/\hat{\rho}_0^\infty$, is equal to that of CMTs (α_ϕ^∞), but its concentration does not depend on stress (due to the normalization of ϕ in Eq. (2.7)). The CMF steady distribution is also independent of the CMT chemical kinetics.

2.5.2 The effect of stress feedback on elastic deformation

Under the previous assumption that stress does not vary much as stiffness evolves in time, the steady-regime elasticity tensor \mathbb{C}_w^∞ is an explicit function of the CMT distribution (Eqs. (2.4) to (2.7), (2.14) and (2.33)). In particular, one can choose (without loss of generality) the surface basis such that \mathbf{e}_x coincides with the main eigenvector of stress. In this basis, the CMF distribution ρ^∞ is even (its odd Fourier coefficients $\{\tilde{\phi}_n^\infty\}_{n \in \mathbb{N}}$

⁶Note that this definition makes sense only for tensile stress ($S_1 \geq S_2 \geq 0$).

⁷This can be mathematically intuited by additionally assuming that $c_0 \gg \int_\pi \phi$, which reduces Eq. (2.9) to a linear ordinary differential equation. Its solution displays an initial slope ($|d_t \phi(\theta, t_0)|$) proportional to $k'_\phi(\mathbf{S}, \theta)$ (that is a decreasing function of f_0).

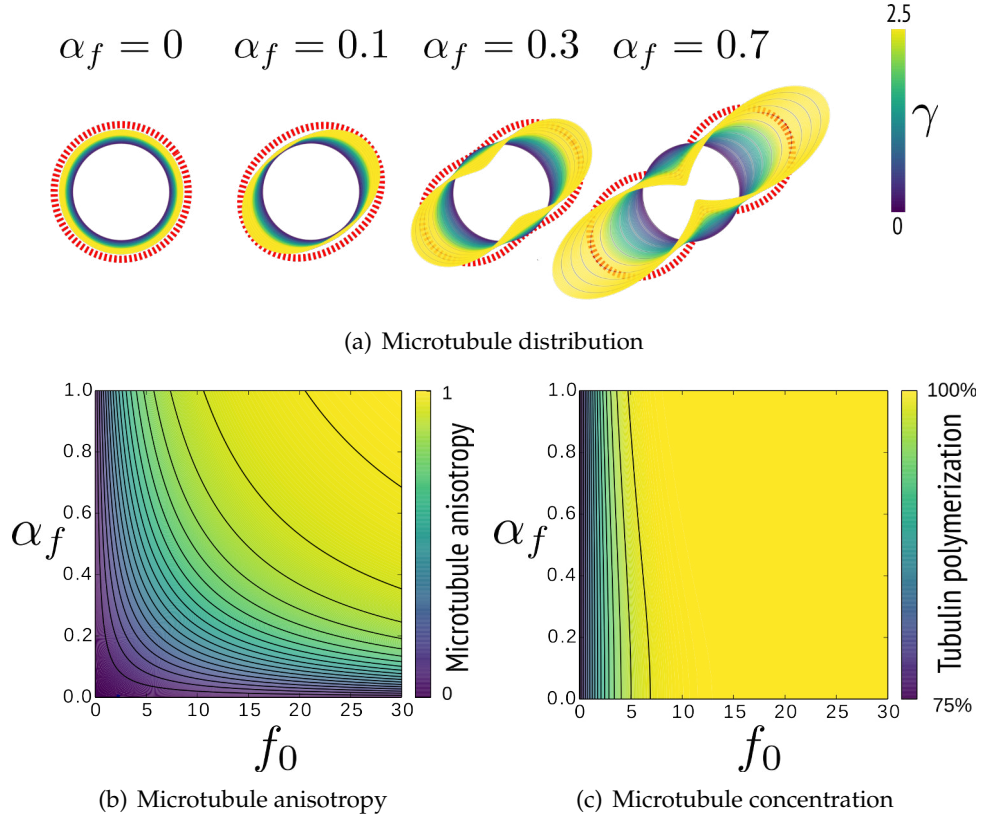


Figure 2.3: **Steady state of the CMT distribution.** Microtubules respond nonlinearly to stress anisotropy *and* amplitude. Microtubule anisotropy depends equally on both stress anisotropy and amplitude. (a) Visualization of the CMT distribution ϕ according to the stress anisotropy α_f ($f_0 = 1$ MPa) and the coupling coefficient γ . The red dashed line represents the polar distribution of forces f . (b) Anisotropy α_ϕ^∞ of the CMT distribution as a function of f_0 (expressed in MPa) and stress anisotropy α_f with stress feedback. (c) Relative concentration of polymerized tubulin $c_p^\infty/c_0 = \pi/2\phi_0^\infty/c_0$ as a function of f_0 (expressed in MPa) and stress anisotropy α_f with stress feedback. Parameters for (b-c): $c_0 = 1 \text{ mol} \cdot \mu\text{m}^{-3}$, $\gamma = 10 \text{ MPa}^{-1}$, $K_\phi = 1$.

vanish) and $[\mathbb{C}_f^\infty]$ becomes:

$$[\mathbb{C}_f^\infty] = \frac{K_\rho Y_f}{8} \begin{bmatrix} 3 + 4\alpha_\phi^\infty + \beta_\phi^\infty & 1 - \beta_\phi^\infty & 0 \\ 1 - \beta_\phi^\infty & 3 - 4\alpha_\phi^\infty + \beta_\phi^\infty & 0 \\ 0 & 0 & 1 - \beta_\phi^\infty \end{bmatrix} \quad (2.15)$$

where $\beta_\phi^\infty = |\hat{\phi}_2^\infty|/\hat{\phi}_0^\infty$ and $K_\rho = k_\rho/k'_\rho$ is the isotropic-stress equilibrium constant of cellulose deposition. By combining the previous equation with the expression of the Fourier coefficients of ϕ (Eq. (2.33)), tensor $\mathbb{C}_w^\infty = \mathbb{C}_f^\infty + \mathbb{C}_g$ can be written as a function of stress, yielding the stress/strain nonlinear relationship illustrated in Fig. 2.4(a). Eq. (2.15) states that the material is asymptotically more resistant to traction in the direction of main stress ($\mathbb{C}_{w,11}^\infty \geq \mathbb{C}_{w,22}^\infty$). Moreover, it is orthotropic (symmetric) according

to the stress axes ($\mathbb{C}_{w,13}^\infty = \mathbb{C}_{w,23}^\infty = 0$)⁸. The expression of stiffness that was obtained is specific to fibrous materials and provides for that more realistic mechanical behavior. This is due to the asymmetry of the CMF distribution, due to a transient phase shift between the respective distributions of CMTs and CMFs, representing the inertia of the system.

In anisotropic elasticity, the respective directions of maximal stress and strain are in general different. In our specific case though, at steady-state, the orthotropy of the elasticity tensor in the stress eigenbasis imposes that both fields (stress and strain) share the same eigenvectors. An alternative remains: either directions of maximal (and minimal) eigenvalues of both fields are parallel or orthogonal. Those two regimes were notably characterized in [Bozorg et al. \(2014\)](#) according to the anisotropy of the material's stiffness. We show how the orthogonality between the main directions of stress and strain can be expressed as a condition on stress only, using the dependence of stiffness on stress. By comparing the ratio

$$\frac{E_\perp^\infty}{E_\parallel^\infty} = \frac{S_2 \mathbb{C}_{w,11}^\infty - S_1 \mathbb{C}_{w,12}^\infty}{S_1 \mathbb{C}_{w,22}^\infty - S_2 \mathbb{C}_{w,12}^\infty} \quad (2.16)$$

(obtained by inverting Eq. (2.2), and illustrated in Fig. 2.4(c)) between longitudinal strain E_\parallel^∞ (collinear to main stress) and orthogonal strain E_\perp^∞ (orthogonal to main stress), to 1, a necessary and sufficient condition for orthogonality can be derived:

$$\alpha_f < \frac{\mathbb{C}_{w,11}^\infty - \mathbb{C}_{w,22}^\infty}{\mathbb{C}_{w,11}^\infty + \mathbb{C}_{w,22}^\infty + 2\mathbb{C}_{w,12}^\infty}. \quad (2.17)$$

By expressing matrix coefficients $\mathbb{C}_{w,ij}^\infty$ according to Eqs. (2.4), (2.5) and (2.15), the previous condition becomes:

$$\alpha_f < \frac{\alpha_\phi^\infty}{1 + 2\eta} \quad \text{with} \quad \eta = \frac{(1 + \nu) Y}{K_\rho Y_f}, \quad (2.18)$$

where η measures the relative stiffness between the gel and the fiber components; the quantity $\alpha_\phi^\infty (1 + 2\eta)^{-1}$ is a measure of the material's anisotropy. This *strain orthogonalization* directly relies on the sharpness of the CMT alignment, and therefore on the stress field (Eqs. (2.11) and (2.35) and Fig. 2.4(c)). Note that since the anisotropy ratio α_ϕ^∞ lies in $[0, 1]$, strain orthogonalization cannot occur for excessive stress anisotropy. In fact, Eq. (2.18) cannot be verified if $\alpha_f \geq (1 + 2\eta)^{-1}$; this limitation comes from the residual isotropy due to the isotropic component (Eq. (2.4)). In brief, we have shown how two mechanical regimes (orthogonality or collinearity of the stress and strain main directions) can be explicitly accounted for by the chemical properties of the CMT array

⁸In the transient regime however, orthotropy cannot be assumed, since the choice of the reference axis does not eliminate both $\tilde{\rho}_1$ and $\tilde{\rho}_2$ in general.

(notably through the sensitivity parameter γ). In particular, *strain orthogonalization* cannot occur if microtubules are less anisotropic than stress (Eq. (2.18)).

Although the number of CMFs, regardless of their orientation, does not depend on stress, we observe differences in the material's areal dilation between the stress-responsive ($\gamma > 0$) and isotropic ($\gamma = 0$) materials. This difference is a general property of fibrous materials, for which areal dilation varies with material anisotropy. Fig. 2.4(b) illustrates the emergence of this effect from stress directly. It allows identifying a particular domain where stress feedback produces higher dilation than in the case of an isotropic, non stress-sensing material. In a biological context, that implies that anisotropic-walled cells could expand faster than isotropic ones, all other parameters being equal. In addition, due to the non-linearity in the strain-based growth equation (Eq. (2.1)), an anisotropic cell may experience one-directional growth, whilst its isotropic counterpart could be kept under the strain threshold in all directions, preventing its irreversible expansion.

2.5.3 Effect of *strain orthogonalization* on tissue-scale development

In order to characterize the impact of the feedback process on an entire structure, we compare the strain field of an inert (non-responsive) material ($\gamma = 0$) and a stress-responsive material ($\gamma > 0$), at steady regime. In fact, Eq. (2.1) formulated in Boudon et al. (2015) expresses the local growth rate as a piece-wise linear function of the strain, suggesting that the elastic response of the tissue directly allows computing its immediate growth. In our context, where no growth is considered, we infer qualitative properties of growth from elastic strain.

One of the simplest shapes, involving anisotropic stresses, is the cylinder. Similar geometries have been studied in the context of plant modeling as they can be considered as idealized stems (Boudon et al., 2015, Bozorg et al., 2014, Hamant et al., 2008, Sassi et al., 2014). We employ here a cylinder closed by a hemispherical cap and fixed at its boundary (Fig. 2.5). In this case, stress is known from continuous mechanics: it is anisotropic in the cylindrical trunk where hoop stress S_1 equals twice axial stress S_2 , and isotropic on the cap (mean stress is equal to S_2), as confirmed by FEM (Figs. 2.5(a) and 2.5(b)).

In absence of feedback ($\gamma = 0$, then CMT distribution is isotropic, Fig. 2.5(c)), circumferential stress results in circumferential strain (Fig. 2.5(e)) that should tend to break the cylindrical shape. By activating the feedback process ($\gamma > 0$), CMTs align in a circumferential way on the trunk, and remain isotropic on the cap since stress provides no directional bias here (Fig. 2.5(d)).

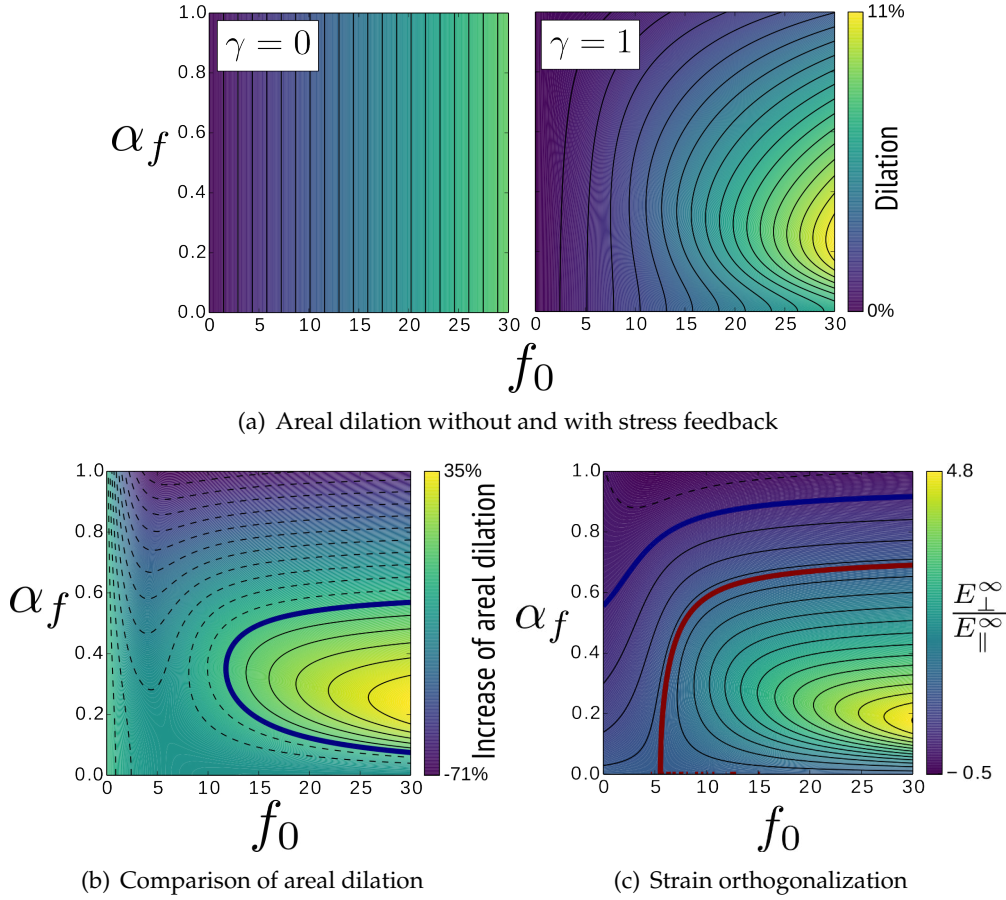


Figure 2.4: The effect of stress feedback on elastic strain. Stress feedback confers non-linear elastic properties to the material. Two noticeable effects are the effect of CMF reorganization on areal expansion (that may differ relative to the isotropic linear case) and the strain and stress respective directions (that may be orthogonal). **(a)** Areal dilation ($\text{Tr } E^{\infty}$) Δ_{OFF} without ($\gamma = 0$) and Δ_{ON} with ($\gamma = 1$, unit: 10^{-6} Pa^{-1}) feedback. In absence of feedback, dilation is a linear function of f_0 , which is not the case with feedback. **(b)** Relative variation $(\Delta_{\text{ON}} - \Delta_{\text{OFF}}) / \Delta_{\text{OFF}}$ (in %) of areal dilation between a stress-responsive material ($\gamma = 1$) and a non-responsive material ($\gamma = 0$), as a function of f_0 (expressed in MPa) and stress anisotropy α_f . Blue bold contour indicates level 0 ($\Delta_{\text{OFF}} = \Delta_{\text{ON}}$). **(c)** Ratio $E_{\perp}^{\infty}/E_{\parallel}^{\infty}$ of strain (in the stress eigenbasis, where the basis vectors are sorted according to the descending order of eigenvalues) as a function of f_0 (expressed in MPa) and α_f , with stress feedback. Main stress and strain are orthogonal *iff* $E_{\perp}^{\infty}/E_{\parallel}^{\infty} > 1$; domains labeled \perp and \parallel respectively indicate those where main stress and strain are orthogonal or collinear. Blue and red bold contours respectively indicate levels 0 (null orthogonal strain $E_{\perp}^{\infty} = 0$) and 1 (isotropic strain $E_{\perp}^{\infty} = E_{\parallel}^{\infty}$). Parameters: c_0 (arbitrary), $\gamma = 10 \text{ MPa}^{-1}$, $K_{\phi} = 1$ (arbitrary), $K_{\rho} = 1 \text{ mol} \cdot \mu\text{m}^{-3}$, $Y_f = 2 \text{ GPa}$, $\nu = 0.2$, $Y = 0.2 \text{ GPa}$.

Enabling the feedback ($\gamma > 0$) may allow strain to become maximal in the axial direction (orthogonal to stress), provided a value of γ satisfying Eq. (2.18) (Figs. 2.5(b) and 2.5(f)). This results in a slimming and an axial stretching of the cylindrical trunk.

One also observes that the dilation of the finite elements (areal expansion, equal to $\text{Tr } E$

for small deformation) is bigger with the feedback than without (Figs. 2.5(e) and 2.5(f)), corresponding to the positive domain in Fig. 2.4(b).

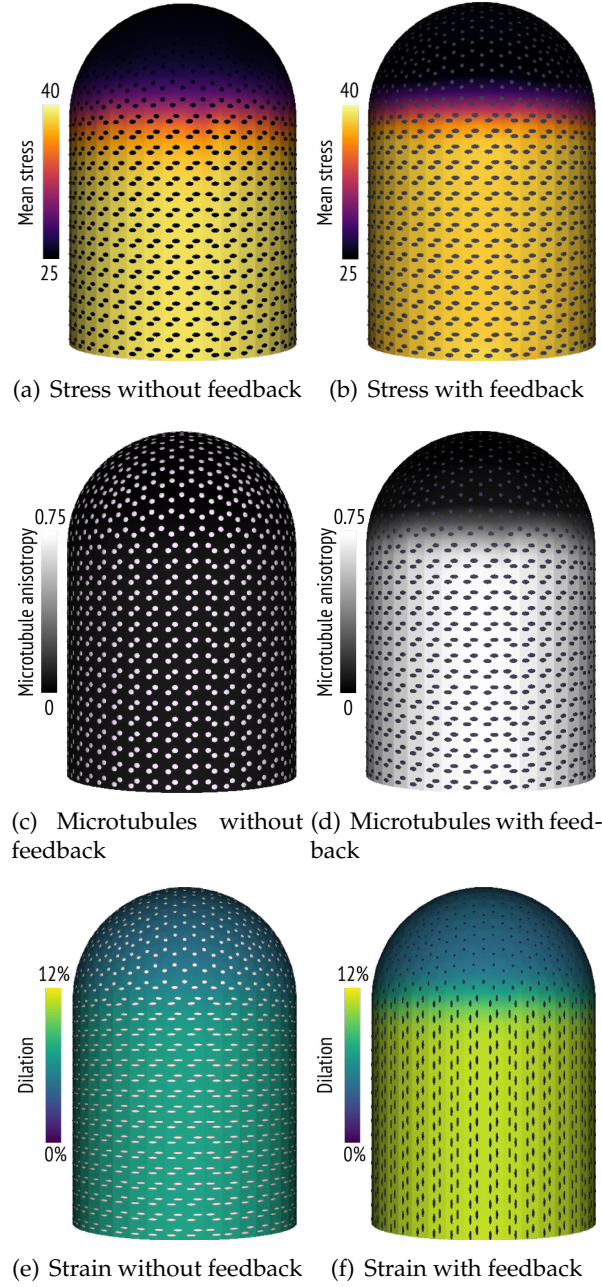


Figure 2.5: **Virtual stem.** Circumferential hoop stresses promote circumferential coalignment of CMTs resulting in axial elongation. **(a-b)** Stress field. Heat map: mean stress $f_0/2$ (in MPa), elliptic representation of the tensor. **(a)** In absence of feedback, **(b)** with feedback. **(c-d)** Microtubule distribution. Heat map: CMT anisotropy α_ϕ^∞ , ellipses: alignment tensor (see Appendix 2.E). **(c)** In absence of feedback, **(d)** with feedback. **(e-f)** Strain field. Heat map: dilation $\text{Tr } \mathbf{E}$. **(e)** In absence of feedback, **(f)** with feedback. Parameters: stem dimensions $300 \mu\text{m} \times 200 \mu\text{m}$, turgor pressure $P = 0.5$ MPa, c_0 (arbitrary), $\gamma = 2 \times 10^{-7} \text{ Pa}^{-1}$, $K_\phi = 1$, $K_\rho = 1 \text{ mol} \cdot \mu\text{m}^{-3}$, $Y_f = 20 \text{ GPa}$, $\nu = 0.2$, $Y = 1 \text{ GPa}$.

The previous simulation involves a positively curved shape. Yet, plants, and in particular the meristem, display several negatively curved zones (for example at the frontier between organs). This scenario has been idealized by merging two sphere meshes, creating a negatively curved junction (Fig. 2.6).

In that case, the outgrowth generates a high circumferential stress at the concave bend between the two spheres, where the gaussian curvature is negative (Figs. 2.6(b) and 2.6(c)). Consequently, the strain field is also highly circumferential in absence of feedback (Fig. 2.6(f)). This contributes to augment the size of the frontier, relative to the reference state.

In contrast, the stress feedback allows to compensate the circumferential stress, and develop axial strains (Figs. 2.6(c) and 2.6(g)). In this case, the decrease of the circumferential strain contributes to maintaining the size of the frontier, reinforcing the separation between the meristem and its primordia, for instance. This example reinforces the idea exposed in [Bozorg et al. \(2014\)](#), that the feedback promotes the anisotropy of shape, preserving non-spherical curvature.

As a supplemental layer of shape complexity, the effect of a topological change has been examined. We represented this scenario through a circular hole in a hemispherical cap (fixed at its boundary), mimicking the ablation experiments described in [Hamant et al. \(2008\)](#), [Sampathkumar et al. \(2014a\)](#).

Here, highly anisotropic stresses are generated at the hole's vicinity (Figs. 2.7(b) and 2.7(c)). In absence of feedback, this would promote an expansion of the hole's size (Fig. 2.7(f)). With feedback, the steady-regime strain features radial orientation (Fig. 2.7(g)) or, at least, less anisotropic circumferential orientation (where strain orthogonalization fails, see Eq. (2.18)), allowing a decrease of the hole's circumference relative to the non-stress-sensing case. This comes with a relaxation of the circumferential stress amplitude (around 10% with the present parameters) at the close periphery of the hole (Figs. 2.7(b) and 2.7(c)). To sum this up, it seems that a strain-based growth, coupled with a stress feedback may tend to close topological defects such as holes.

2.6 Discussion

Existing modeling efforts assessing the feedback mechanism between mechanics and growth in plant tissues rely on a phenomenological *black box* to express the influence of stress on cell wall stiffness ([Bozorg et al., 2014](#), [Hamant et al., 2008](#), [Hervieux et al., 2016](#)). This work is an effort to open this black box by making a first step toward a more quantitative, mechanistic, multiscale analysis of stress-based feedback. This approach

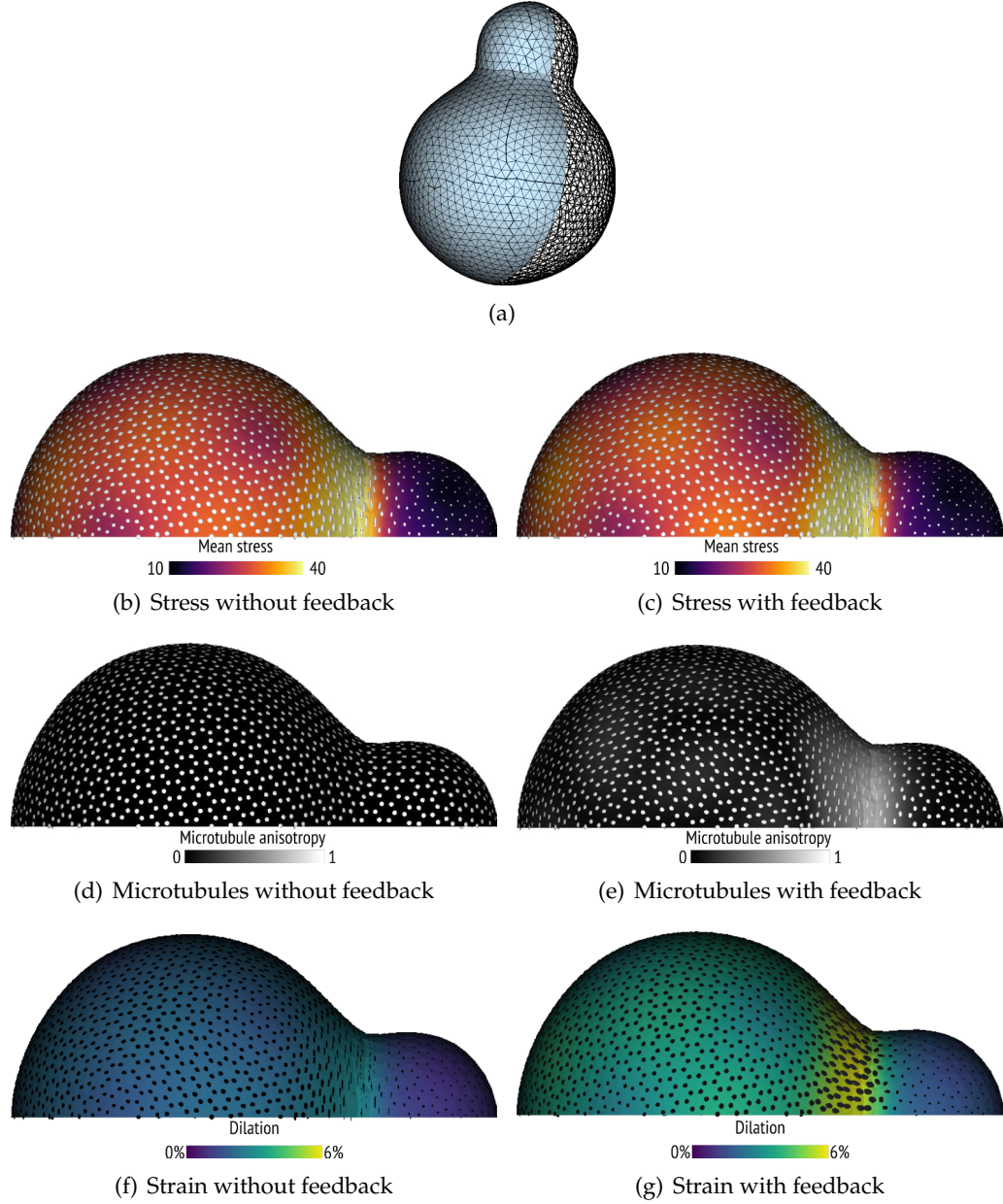


Figure 2.6: **Virtual outgrowing organ.** Highly anisotropic stresses at the concave frontier contribute to maintain the latter that expands mainly axially. (a) Original mesh and visualized cut (in blue). (b-c) Stress field. Heat map: mean stress $f_0/2$ (in MPa), elliptic representation of the tensor. (b) In absence of feedback, (c) with feedback. (d-e) Microtubule distribution. Heat map: CMT anisotropy α_ϕ^∞ , ellipses: alignment tensor (see Appendix 2.E). (d) In absence of feedback, (e) with feedback. (f-g) Strain field. Heat map: dilation $\text{Tr } \mathbf{E}$. (f) In absence of feedback, (g) with feedback. Parameters: mesh characteristic dimension $100 \mu\text{m}$ (order of magnitude), turgor pressure $P = 0.5 \text{ MPa}$, c_0 (arbitrary), $\gamma = 10 \text{ MPa}^{-1}$, $K_\phi = 1$, $K_\rho = 1 \text{ mol} \cdot \mu\text{m}^{-3}$, $Y_f = 20 \text{ GPa}$, $\nu = 0.2$, $Y = 0.3 \text{ GPa}$.

relies on a parsimonious description of the chemical behavior of cortical microtubules, coupled to cellulose microfibril deposition. A homogenization procedure allows us to derive from the angular distribution of these microfibrils a continuous expression

of stiffness as a function of its low-frequency Fourier coefficients, as described in Cox (1952). In this formalization, the material stiffening in the direction of main tension results from the anisotropization of the microtubule array due to the stress field. The coupling between a linear constitutive relation (Hooke's law, Eq. (2.2)) and a nonlinear dependency of the elastic properties upon stress yields an integrated formulation of the behavior, in which stiffness is expressed as a function of stress itself:

$$\mathbf{E}^\infty = \mathbb{C}_w^\infty (\mathbf{S})^{-1} : \mathbf{S}. \quad (2.19)$$

This expression captures the nonlinear behavior of the wall depicted in Figs. 2.3 and 2.4.

Although growth is not included, the model can be useful in exploring the role of a stress-based feed-back on growth. Indeed, assuming that growth is a piece-wise linear function of strain (Eq. (2.1), Boudon et al., 2015, Bozorg et al., 2016), the predicted strain patterns can be used as a proxy for the immediate growth.

In a situation where elasticity is isotropic or slightly anisotropic, strain and stress share the same main axis. Our FEM simulations suggest that in that case strain-based growth should favor spherical shapes, while cylinders or more complex shapes cannot be obtained (see also Bozorg et al., 2014). Indeed, these shapes require that the main strain and stress directions are orthogonal. This is possible only if the elastic anisotropy of the material is higher than the stress anisotropy (Ineq. (2.18)). *In vivo*, expansion can occur in parallel or perpendicular to the maximal stress direction, depending on the type of tissue (Burian et al., 2013). Our model captures this complexity thanks to the exponential form of the microtubule degradation rate (Eq. (2.10)). Indeed, this superlinear amplification of the differential of force in the microtubule distribution (Fig. 2.3(a)), allows the system to satisfy the *strain orthogonalization* condition (Ineq. (2.18)) in specific cases, such as along an axis or around a hole or a bump. Fig. 2.4(c) shows that orthogonalization may happen, provided a sufficient stress amplitude f_0 , but fails with excessive stress anisotropy (2.18), Fig. 2.4(c)) due to the presence of an isotropic matrix. The regulation of the matrix through the dimensionless ratio η , should affect the mean stiffness and anisotropy of the material. Such a regulation, e.g. occurring through auxin-induced alteration of the pectin gel, has been experimentally reported (Braybrook and Peaucelle, 2013, Peaucelle et al., 2011).

Based on the strain orthogonalization condition (Ineq. (2.18)), our model allows to make predictions on the growth behavior in different biological scenarios. In the case of wound-induced stress, idealized in our simulations conducted on a punctured dome (Fig. 2.5, Hamant et al., 2008, Sampathkumar et al., 2014a), stiffening in the circumferential direction leads to maximal strain in the radial direction. Supposedly, this could enhance the efficiency of a wound healing process based on strain-based growth.

Similarly, our simulations predict that a bulging primordium generates a highly anisotropic stress pattern around itself. This, in turn, should sharply amplify growth heterogeneity between the outgrowth and its surroundings (Fig. 2.6). More generally, in the previously studied scenarios, stress feedback tends to modify the strain field, counteracting the natural tendency of pressure to induce a spherical shape, preserving the negative curvature at the periphery of an organ or promoting axial expansion (Fig. 2.5).

In addition, our model includes crucial molecular processes underlying the control of wall mechanics. The degradation term expressed by Eq. (2.10) can, *in vivo*, be related to *katanin*-based microtubule severing. Katanin is a major regulator of microtubule dynamics in both plants and animals and is involved in the formation of anisotropic microtubular arrays (Burk and Ye, 2002). Although the precise mechanism is not understood, it facilitates microtubule ordering along the main stress direction (Uyttewaal et al., 2012). At a molecular scale, the stabilization of CMTs by stress can be related to possible stress-induced conformational changes undergone by tubulin (Landrein and Hamant, 2013a, Sampathkumar et al., 2014a), analogous to those occurring in animal cortical biopolymers (Ghanti et al., 2016). In the context of an anisotropic stress field, the tensile forces exerted on microtubules and, therefore, the efficiency of the katanin activity vary from one direction to another. We modeled this molecular mechanism as a stress-dependent expression of the microtubule depolymerization rate (Eq. (2.10)). This expression, inspired by studies on other force-sensitive biomolecular systems (Bell, 1978, Changeux, 2012, Nicolas et al., 2004), depicts the idea that the force-induced conformational change of microtubules modifies the enthalpy of the katanin-microtubule reaction. More specifically, its activation barrier is increased by the work of tensile forces exerted on microtubules, leading to their stabilization. The resulting exponential dependency of the degradation rate on tensile force enables the amplification of the microtubule response to mechanical loading. For the sake of simplicity, we neglected interactions between microtubules in the dynamics described by Eq. (2.9). In a finer model, those should be integrated as well, for they support the capacity of self-alignment of microtubule arrays (Allard et al., 2010, Dixit and Cyr, 2004, Tindemans et al., 2010) and could enhance the stress-sensitive mechanism we describe here.

Experimental tools to quantify microtubule distributions in plant tissues have been developed recently (Boudaoud et al., 2014, Tsugawa et al., 2016). Combining these measurement techniques with our modeling approach, applied to high-quality digitalized images (Cerutti et al., 2017) would be a way to challenge some of the results exposed here, such as, the dependency of microtubule anisotropy α_ϕ on the mean stress f_0 . It could also help estimate the transduction coefficient γ which plays a central role in the sensitivity of the feedback mechanism. This could constitute a fruitful line of future work.

Finally, our model raises further questions about the dynamics of microfibrils. The latter is described by Eq. (2.7) that contains a degradation term. This term is required mathematically for the existence of a steady regime. Without it, the cell wall would undergo ceaseless accumulation of cellulose. Whether such a limiting term has an enzymatic origin or could be explained by a growth-related dilution/advection process remains to be established. It is in fact not self-evident that growth-induced advection/diffusion could suffice for rapid changes in growth direction (as observed during organ initiation). Indeed, in principle, newly deposited microfibrils would be in mechanical competition with older layers with different orientation. This competition would further slow the mechanical responsiveness of the tissue down. In depth theoretical modeling should provide further insights into this problem.

2.7 Acknowledgments

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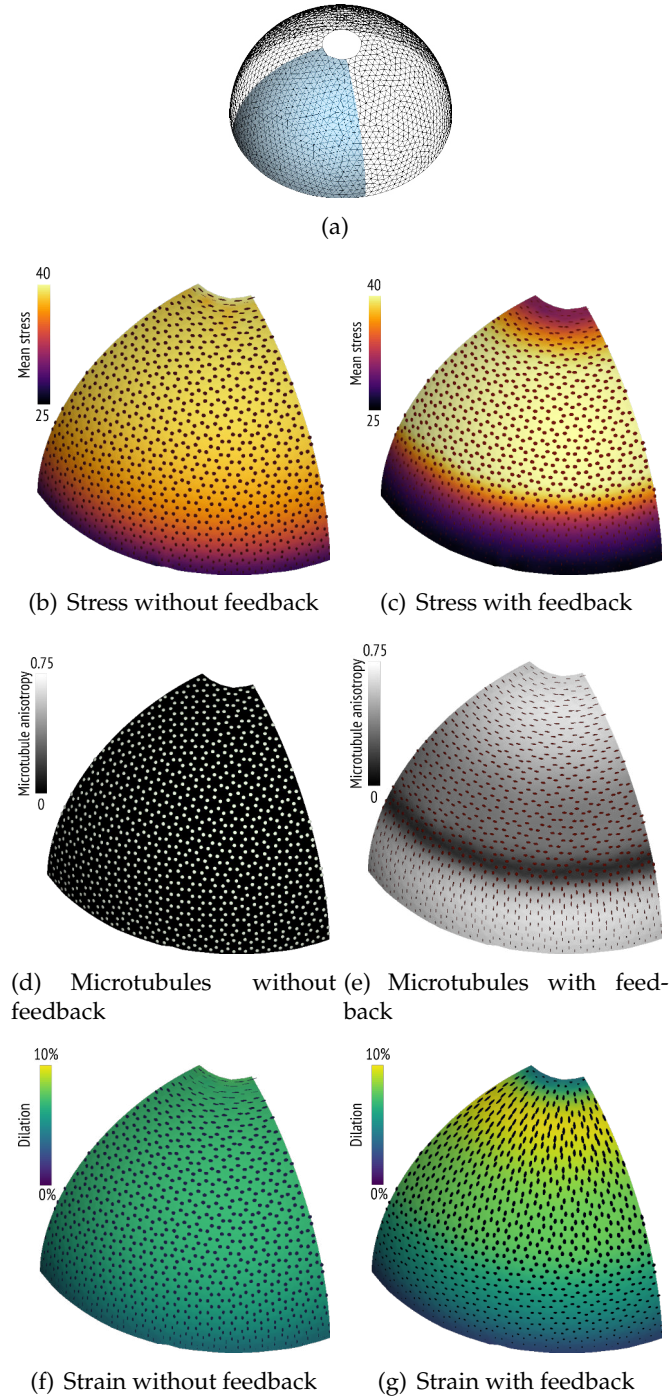


Figure 2.7: **Holed cap.** Circular reorganization of CMTs allows to limit the effect of the breach by promoting radial expansion around the hole. (a) Original mesh and visualized cut (in blue). (b-c) Stress field. Heat map: mean stress $f_0/2$ (in MPa), elliptic representation of the tensor. (b) In absence of feedback, (c) with feedback. (d-e) Microtubule distribution. Heat map: CMT anisotropy α_ϕ^∞ , ellipses: alignment tensor (see Appendix 2.E). (d) In absence of feedback, (e) with feedback. (f-g) Strain field. Heat map: dilation $\text{Tr } \mathbf{E}$. (f) In absence of feedback, (g) with feedback. Parameters: mesh characteristic dimension $100 \mu\text{m}$ (order of magnitude), turgor pressure $P = 0.5 \text{ MPa}$, $\gamma = 2 \times 10^{-7} \text{ Pa}^{-1}$, c_0 (arbitrary), $K_\phi = 1$, $K_\rho = 1 \text{ mol} \cdot \mu\text{m}^{-3}$, $Y_f = 20 \text{ GPa}$, $\nu = 0.2$, $Y = 0.5 \text{ GPa}$.

Appendices

2.A The elasticity tensor as a function of the microfibril distribution

This appendix details Eq. (2.6), that establishes a relation between the fiber organization and the associated stiffness, as described in [Cox \(1952\)](#).

We consider that CMFs behave like linear springs (of rest length l_0 and spring constant k). Moreover, by assuming affine deformation, their deformation reads:

$$\frac{\Delta l(\theta)}{l_0} \simeq \mathbf{e}_\theta \cdot (\mathbf{E} \cdot \mathbf{e}_\theta) = \mathbf{E} : \mathbf{e}_\theta \otimes \mathbf{e}_\theta = \mathbf{E} : \mathbf{\Theta}.$$

where $\mathbf{\Theta} = \mathbf{e}_\theta \otimes \mathbf{e}_\theta$ is the projector on direction θ (' \otimes ' depicts the tensor outer product). The previous expression yields the total stretching energy density per unit volume of fibers (summed over all directions):

$$\Psi = \frac{1}{2} Y_f \int_{\pi} d\theta \rho(\theta) (\mathbf{E} : \mathbf{\Theta})^2 = \frac{1}{2} \mathbf{E} : Y_f \int_{\pi} d\theta \rho(\theta) \mathbf{\Theta} \otimes \mathbf{\Theta} : \mathbf{E} \quad (2.20)$$

with:

$$Y_f = \frac{k l_0^2}{\epsilon}$$

(where ϵ depicts wall thickness). Deriving twice the energy yields the elasticity tensor:

$$\mathbb{C}_f = \frac{\partial^2 \Psi}{\partial^2 \mathbf{E}} = Y_f \int_{\pi} d\theta \rho(\theta) \mathbf{\Theta} \otimes \mathbf{\Theta}$$

or in the basis $\{\mathbf{e}_x, \mathbf{e}_y\}$:

$$(\mathbb{C}_f)_{ijkl} = Y_f \int_{\pi} d\theta \rho(\theta) \cos(\theta)^{\Delta_{ijkl}} \sin(\theta)^{4-\Delta_{ijkl}}$$

with $\Delta_{ijkl} = \delta_{i,1} + \delta_{j,1} + \delta_{k,1} + \delta_{l,1}$, where ' δ ' stands for the Kronecker delta function. Linearizing the cosine-sine products yields an expression of \mathbb{C}_f involving the 0th, 1st

and 2nd order Fourier coefficients:

$$[\mathbb{C}_f] = \frac{\pi Y_f \rho_0}{16} \begin{bmatrix} 3 + \frac{\rho_2 + 4\rho_1}{\rho_0} & 1 - \frac{\rho_2}{\rho_0} & \frac{2\tilde{\rho}_1 + \tilde{\rho}_2}{\rho_0} \\ 1 - \frac{\rho_2}{\rho_0} & 3 + \frac{\rho_2 - 4\rho_1}{\rho_0} & \frac{2\tilde{\rho}_1 - \tilde{\rho}_2}{\rho_0} \\ \frac{2\tilde{\rho}_1 + \tilde{\rho}_2}{\rho_0} & \frac{2\tilde{\rho}_1 - \tilde{\rho}_2}{\rho_0} & 1 - \frac{\rho_2}{\rho_0} \end{bmatrix} \quad (2.21)$$

where, for any natural number n :

$$\hat{\rho}_n = \int_{\pi} \frac{d\theta}{\pi} \rho(\theta) e^{-2in\theta} \quad \text{and} \quad (\rho_n, \tilde{\rho}_n) = 2(\Re(\hat{\rho}_n), -\Im(\hat{\rho}_n)). \quad (2.22)$$

2.B Modeling cellulose deposition via CSC trajectories

We here develop a kinetic model for CSC-CMT binding, providing a chemical background to the cellulose deposition model. Here, cellulose deposition is controlled through the orientation of the CSC trajectories depicted by an *a priori* non-uniform π -periodic distribution $\mu(\theta)$ that depicts the trajectories of proteins (regardless of their sense).

$$d_t \rho(\theta) = k_{\rho}^0 \mu(\theta) - k'_{\rho} \rho(\theta), \quad (2.23)$$

We assume that a constant pool of CSCs is available, characterized by a constant concentration μ_{tot} . Proteins can be either free or bound to a CMT. The free proteins are assumed to follow individual Brownian trajectories within the membrane bi-layer; for that the related cellulose deposition is isotropic, characterized by a concentration μ_{free} . The contribution of the bound proteins is characterized by the distribution $\mu_{\text{bound}}(\theta)$ that depicts the trajectories of proteins on CMTs. We assume that CSCs are active regardless of their binding state, implying that cellulose deposition occurs even in the absence of CMTs (Emons et al., 2007). Microtubules solely introduce a bias in the distribution of trajectories μ . The total concentration of CSCs reads:

$$\int_{\pi} \mu = \mu_{\text{free}} + \int_{\pi} \mu_{\text{bound}} = \mu_{\text{tot}}. \quad (2.24)$$

The kinetics of CSC-CMT binding is modeled by a multiplicative coupling between the concentration of CMTs in a given direction θ and the concentration of available free proteins, along with an isotropic liberation rate:

$$d_t \mu_{\text{bound}}(\theta) = k_{\mu} \mu_{\text{free}} \phi^{\infty}(S, \theta) - k'_{\mu} \mu_{\text{bound}}(\theta) \quad (2.25)$$

where k_μ and k'_μ are kinetic constants. Eqs. (2.24) and (2.25) lead to the steady-state expression:

$$\mu_{\text{bound}}^\infty(\theta) = \mu_{\text{tot}} \cdot \frac{K_\mu \phi^\infty(\mathbf{S}, \theta)}{1 + K_\mu \int_\pi \phi^\infty} \quad (2.26)$$

where $K_\mu = k_\mu/k'_\mu$ depicts the affinity of CSCs for CMTs. By combining the previous steady-state distribution of CSCs with the expression of cellulose deposition (Eq. (2.23)), we obtain:

$$d_t \rho(\theta) = k_\rho^0 \mu_{\text{tot}} \cdot \frac{1 + K_\mu \phi^\infty(\mathbf{S}, \theta)}{1 + K_\mu \int_\pi \phi^\infty} - k'_\rho \rho(\theta) \quad (2.27)$$

that can be roughly simplified to obtain Eq. (2.7) by assuming that for all θ , $K_\mu \phi^\infty(\mathbf{S}, \theta) \gg 1$ (strong affinity):

$$d_t \rho(\theta) = k_\rho^0 \mu_{\text{tot}} \cdot \frac{\phi^\infty(\mathbf{S}, \theta)}{\int_\pi \phi^\infty} - k'_\rho \rho(\theta). \quad (2.28)$$

The product $k_\rho^0 \mu_{\text{tot}}$ defines the kinetic constant k_ρ used in the main text.

2.C Details about the specific expression of the CMT depolymerization probability

Eq. (2.10) derives from Arrhenius law: a chemical reaction rate (k_A hereafter) is a direct function of its activation energy, *i.e.* the energetic barrier (E_A) the system has to overcome to go from one chemical state to another:

$$k_A = k_0 e^{-\frac{E_A}{RT}}. \quad (2.29)$$

The product RT corresponds to the thermal energy of the considered molecules are embedded in (R and T respectively stand for the Arrhenius constant and the absolute temperature).

This general experimental law has been amended latter on in the context of biochemistry and cell adhesion in the following way (Bell, 1978). The energy barrier molecules have to overcome can be lowered or raised when mechanical forces are applied to them. The resulting expression reads:

$$k_B = k_0 e^{-\frac{E_A \pm \alpha f}{RT}} \quad (2.30)$$

where f depicts the force applied to the considered molecules and $\pm \alpha$ stands as a coupling parameter. Clearly, the sign of this coupling parameter depends on the action of the force: if the latter eases the reaction, then the coupling parameter must be negative, so the activation barrier is lowered and the transition rate is increased.

Our working hypothesis is that mechanical forces applied to microtubules reduce their depolymerization rate; we therefore postulated a positive coupling parameters. Simplifying the notation as follows:

$$\begin{cases} k'_\phi \stackrel{\text{def.}}{=} k_0 e^{-\frac{E_A}{RT}} \\ \gamma \stackrel{\text{def.}}{=} \frac{\alpha}{RT} \end{cases} \quad (2.31)$$

and considering that E_A is proportional to force, leads to Eq. (2.10).

2.D Microtubule principal axis and anisotropy at steady regime

Generally, the CMT distribution ϕ is summarized through the angle of its mean orientation (θ_ϕ) and through its anisotropy (α_ϕ), respectively given by:

$$\theta_\phi = -\frac{\arg \hat{\phi}_1}{2} \quad \text{and} \quad \alpha_\phi = \frac{|\hat{\phi}_1|}{\hat{\phi}_0} \quad (2.32)$$

where 'arg' is the complex argument, and $\{\hat{\phi}_n\}_{n \in \mathbb{N}}$ depict the Fourier coefficients of ϕ . We define θ_ρ and α_ρ in the same way, using the CMF distribution ρ .

The steady-state Fourier sequence of CMTs is derived from Eq. (2.11) and read:

$$\forall n \in \mathbb{N} : \quad \hat{\phi}_n^\infty = \frac{c_0}{\pi} \frac{I_n(2\gamma|\hat{f}_1|)}{\frac{e^{-\gamma f_0}}{\pi K_\phi} + I_0(2\gamma|\hat{f}_1|)} e^{-2in\theta_S} \quad (2.33)$$

where θ_S depicts the angle of main stress, and where functions $\{I_n\}_{n \in \mathbb{N}}$ depict the modified Bessel functions of the first kind ([Abramowitz and Stegun, 1972](#)):

$$\forall n \in \mathbb{N}, \forall x \in \mathbb{R} : I_n(x) = \int_{-\pi}^{\pi} \frac{d\theta}{\pi} e^{x \cos \theta} \cos n\theta. \quad (2.34)$$

By expressing the complex argument of $\hat{\phi}_1$ from the previous expression, one sees that $\theta_\phi^\infty = \theta_S$. This means that CMTs align with the main direction of stress. This alignment is moreover symmetric, according to this direction (with the parametrization $\theta \leftarrow \theta - \theta_S$, the imaginary part of the Fourier coefficients vanishes).

The anisotropy ratio α_ϕ^∞ (Fig. 2.3(b)) reads:

$$\alpha_\phi^\infty = \frac{I_1(2\gamma|\hat{f}_1|)}{I_0(2\gamma|\hat{f}_1|)}. \quad (2.35)$$

which is a growing function of $|\hat{f}_1|$ (if $\gamma > 0$). According to the previous expression, the steady-state anisotropy of CMTs depends on stress only, and not on the chemical kinetics.

2.E Details on the simulation pipeline

We detail the numerical procedure employed in Section 2.5.3. All simulations have been performed in the framework presented in (Boudon et al., 2015), based on the open-source softwares *Sofa* (Faure et al., 2012) and *OpenAlea* (Pradal et al., 2008). The numerical procedure employed in this paper is adapted from Boudon et al. (2015).

2.E.1 Mechanical equilibrium

2.E.1.1 Space discretization

Finite element procedure We discretize the continuous model through triangular finite elements. Current material positions \mathbf{x} are expressed according to the mesh nodes $\mathbf{q}^{(n)}$ based on a Lagrange \mathbb{P}_1 barycentric interpolation:

$$\mathbf{x} = \sum_{n \in \mathcal{N}} N^{(n)}(\mathbf{X}) \mathbf{q}^{(n)} \quad (2.36)$$

where \mathcal{N} depicts the set of mesh nodes; $N^{(n)}(\mathbf{X})$ depicts the barycentric shape function associated with node n and evaluated at material position \mathbf{X} . Spatial differentiation yields one 3×2 barycentric deformation gradient tensors per triangle τ :

$$\mathbf{F}^{(\tau)} = \sum_{n \in \mathcal{N}} \mathbf{q}^{(n)} \otimes \frac{\partial N^{(n)}}{\partial \mathbf{X}} \Big|_{\tau} \quad (2.37)$$

The 2D Green-Lagrangian strain tensor and the second Piola-Kirchhoff stress are derived from $\mathbf{F}^{(\tau)}$ according to:

$$\mathbf{E}^{(\tau)} = \frac{1}{2} \left(\left(\mathbf{F}^{(\tau)} \right)^T \cdot \mathbf{F}^{(\tau)} - \mathbf{I} \right) \quad \text{and} \quad \mathbf{S}^{(\tau)} = \mathbb{C}_w^{(\tau)} : \mathbf{E}^{(\tau)}, \quad (2.38)$$

yielding the internal energy of the system (that is only due to deformation):

$$U = \sum_{\tau \in \mathcal{T}} \epsilon \mathcal{A}^{(\tau)} \mathbf{E}^{(\tau)} : \mathbf{S}^{(\tau)} = \sum_{\tau \in \mathcal{T}} \frac{\epsilon \mathcal{A}^{(\tau)}}{2} \mathbf{E}^{(\tau)} : \mathbb{C}_w^{(\tau)} : \mathbf{E}^{(\tau)}, \quad (2.39)$$

where ϵ and $\mathcal{A}^{(\tau)}$ respectively stand for the wall thickness (that is uniform) and the surface area (reference configuration) of triangle τ . Nodal inner forces are obtained from the gradient of the strain energy with respect to node positions:

$$\mathbf{f}_{\text{int}}^{(n)} = -\frac{\partial U}{\partial \mathbf{q}^{(n)}}. \quad (2.40)$$

The external loads are derived from pressure and expressed at each node n according to its adjacent triangles \mathcal{T}_n :

$$\mathbf{f}_{\text{ext}}^{(n)} = \frac{P}{3} \sum_{\tau \in \mathcal{T}_n} a^{(\tau)} \mathbf{n}^{(\tau)}, \quad (2.41)$$

where $\mathbf{n}^{(\tau)}$ stands for the outer normal of triangle τ ; $a^{(\tau)}$ is the surface area of triangle τ in the current configuration.

Meshes All meshes have been generated through the open source software *Blender* (blender.org). At the exception of the cylindrical stem (for which the original mesh was satisfactory), they have been enhanced via a surface smoothing procedure implemented in the non-commercial meshing tool *Graphite* (alice.loria.fr/software/graphite).

2.E.1.2 Time discretization

Computing the equilibrium (Eq. (2.3)) consists in minimizing the total energy U (see [Boudon et al., 2015](#), for further details). This is performed by solving the equation of motion until node forces balance. We use the backward Euler method and the conjugate gradient to perform this step.

To improve stability and to avoid excessive stiffness inhomogeneity, the tensor \mathbf{S} employed in Eq. 2.11 is modified on each element and integrates a weighted average of the neighboring stress tensors.

2.E.2 Dynamical resolution of Eq. 2.7

The dynamics of the CMF distribution is computed by an iterative update of the first three Fourier coefficients of ρ through the following forward Euler time scheme with constant time step Δt (Eq. 2.7 and Fig. 2.8):

$$\forall n : \quad \hat{\rho}_n^{t+\Delta t} = (1 - k'_\rho \Delta t) \hat{\rho}_n^t + \frac{2k_\rho \Delta t}{\pi} \frac{\hat{\phi}_n^\infty(\mathbf{S}^t)}{\hat{\phi}_0^\infty(\mathbf{S}^t)}. \quad (2.42)$$

The stiffness tensor \mathbb{C}_w is subsequently updated according to Eqs. 2.4 to 2.6 and 2.33.

At $t = t_0$, the system is set to its null-stress equilibrium, derived from from Eqs. 2.11 and 2.14:

$$\phi(\theta, t_0) = \frac{c_0 K_\phi}{1 + K_\phi \pi} \quad \text{and} \quad \rho(\theta, t_0) = \frac{K_\rho}{\pi}. \quad (2.43)$$

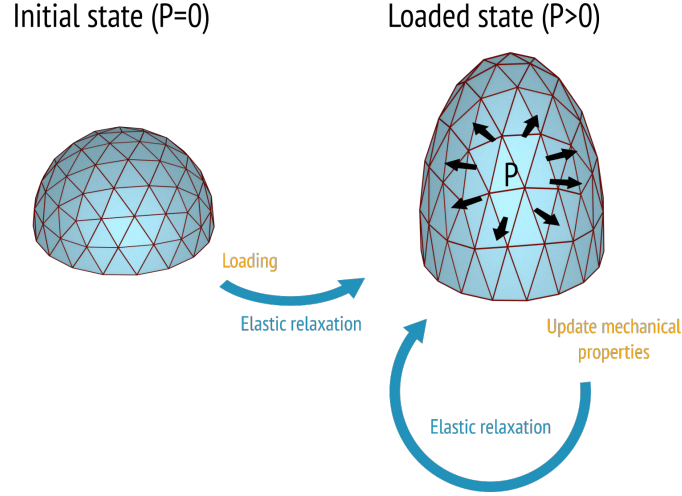


Figure 2.8: **Simulation pipeline.** The initial state corresponds to the null stress state (no residual stress and no loading). The elastic equilibrium is computed after loading the structure with pressure $P > 0$. At each elastic equilibrium, the mechanical properties are updated, which brings the systems out of equilibrium.

2.E.3 Visualization of the CMF distributions in Figs. 2.5 to 2.7

We adapted the concept of nematic tensor from the physics of liquid crystals (De Gennes and Prost, 1995) by defining:

$$\mathbf{Q}_\phi = 4 \int_\pi \frac{d\theta}{\pi} \phi(\theta) \boldsymbol{\Theta} = \phi_0 \begin{bmatrix} 1 + \frac{\phi_1}{\phi_0} & \frac{\tilde{\phi}_1}{\phi_0} \\ \frac{\tilde{\phi}_1}{\phi_0} & 1 - \frac{\phi_1}{\phi_0} \end{bmatrix}. \quad (2.44)$$

Notice that, as stated in 2.5.1, the previous definition builds a formal equivalence between the definition of anisotropy based on Fourier coefficients (Eqs. (2.13) and (2.32)) and the nematic-based ratio employed in Boudaoud et al. (2014) (Eq. (2.12)). In fact, the matrix expression of \mathbf{Q}_ϕ becomes diagonal in the basis associated with the rotation that zeroes the odd Fourier coefficient $\tilde{\phi}_1$ (in this basis $|\hat{\phi}_1| = \hat{\phi}_1 = \phi_1/2 \in \mathbb{R}^+$). Since $|\hat{\phi}_1|$ and ϕ_0 are base-invariant, the two eigenvalues of \mathbf{Q}_ϕ are equal to $\phi_0 \pm 2|\hat{\phi}_1|$, which gives:

$$\frac{Q_{\phi,1} - Q_{\phi,2}}{Q_{\phi,1} + Q_{\phi,2}} = \frac{2|\hat{\phi}_1|}{\phi_0} = \frac{|\hat{\phi}_1|}{\hat{\phi}_0} \quad (2.45)$$

\mathbf{Q}_ϕ is equivalent to an ellipse with main axis oriented by $\theta_\phi = -1/2 \arg \hat{\phi}_1$ (Eq. 2.32), and with aspect ratio and semi-major axis respectively given by $(1 - \alpha_\phi) / (1 + \alpha_\phi)$ (with $\alpha_\phi = |\hat{\phi}_1|/\hat{\phi}_0$) and $\phi_0 (1 + \alpha_\phi)$ (see Fig. 2.9).

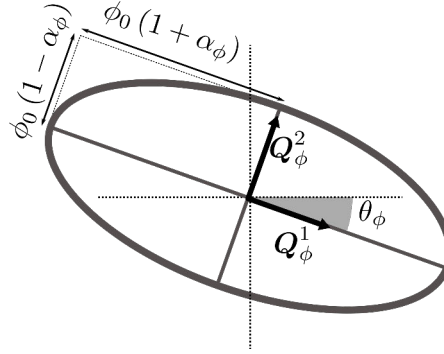


Figure 2.9: Ellipse-based visualization of the nematic tensor associated with the distribution ϕ .

Chapter 3

Coupling growth and stress feedback

3.1 Introduction

In the previous chapter, we developed a model of the stress feedback valid in the elastic regime (namely neglecting growth). In this chapter, we detail a method to couple the previous model to growth, using the growth model developed in [Boudon et al. \(2015\)](#). By doing so, we model the complete feedback loop that was introduced in Chapter 1 and illustrated in Fig. 1.11). For this coupling to be physically consistent, we model the *advection*-like deformation undergone by the fiber arrays as the infinitesimal volumes of the tissue deform.

Here, we will study the dynamical properties of the system (stress feedback + growth), in particular, in the context of symmetry¹ breaking and symmetry maintenance, by using ideally-shaped pressurized shells. Indeed, the fact that the feedback exploits stress anisotropy to generate anisotropic growth suggests that it has the potential to break an initial symmetry of shape by responding to slight geometrical asymmetries. To study this effect, we here consider the highest level of symmetry, that is the spherical shape, and we assess its stability under various kinds of feedback parameters and shape perturbations. We explore the properties of the model focusing on two main key parameters, *i.e.* the sensitivity and rapidity of the feedback (detailed hereafter), that need to be modulated in order to promote efficient asymmetry amplification. From a biological

¹The notion of symmetry is here to be understood in the sense used in physics, that is the existence of a transformation that leaves some property – here the shape – of an object unchanged.

standpoint, this kind of process may be involved in the early stages of organ development, for example allowing for small roundish bumps to grow towards axisymmetric, highly elongated organs like branches.

3.2 Model

3.2.1 Modeling growth and growth-induced fiber advection

Growth is modeled as in [Boudon et al. \(2015\)](#). We consider a morphoelastic material that undergoes rapid elastic relaxation and slow plastic expansion (see Section 1.2.2), both combined through the multiplicative decomposition of the deformation described by Eq. (1.6). Inasmuch as the elastic and plastic regimes occur with very different time scales, one can simulate the incremental evolution of the system's geometry through a two-timing procedure, consisting of i/ computing the elastic equilibrium of the system and ii/ modifying the reference of the material.

The elastic regime of the material is already described in Chapter 2, in which we have detailed a constitutive relation for the material (see Eqs. (2.4) to (2.7) and (2.11)) taking into account stress-induced CMT reorganization and CMF turnover. As for the plastic regime, we use the strain-based growth expressed by Eq. (1.12) ([Boudon et al., 2015](#)), and discretized by the following forward Euler procedure:

$$\mathbf{F}_g(t + \Delta t) = (\mathbf{I} + \Delta t \mathbf{L}_g(t)) \cdot \mathbf{F}_g(t) \quad (3.1)$$

where $\mathbf{L}_g(t)$ depends on the Green-Lagrangian strain $\mathbf{E}(t) \stackrel{\text{def.}}{=} \frac{1}{2} (\mathbf{F}_e^T(t) \cdot \mathbf{F}_e(t) - \mathbf{I})$ at step t as expressed by Eq. (1.12); Δt is a constant time step size. The factor $\mathbf{f}_g(t) \stackrel{\text{def.}}{=} \mathbf{I} + \Delta t \mathbf{L}_g(t)$ depicts the incremental growth that is applied at each growth time step.

In principle, growth induces a change in the fiber density and orientation, which can be assimilated to an advective process. In fact, by virtue of mass conservation, the volumetric density of CMFs decreases as the wall expands if we assume that no cellulose intake is involved. Moreover, anisotropic growth modifies the orientation each fiber, which rotates towards the direction of main deformation.

To model these effects, we assume that the deformation of the fibers is *affine*, i.e. that these deform according to the macroscopic medium (they neither rotate nor slide relative to the material). The growth procedure detailed previously (Eq. (3.1)) naturally provides an incremental way to simulate the advection of the CMFs. At each growth step, the medium is updated with a small incremental deformation \mathbf{f}_g^n that affects the fiber density function (*N.B.*: to simplify notations, we drop the exponent n in the rest

of this paragraph, since all the next developments are made at time step n). Let $R(\Theta)$ and $\rho(\theta)$ be the angular density of CMFs (in the reference configuration) respectively before and after applying the incremental deformation \mathbf{f}_g . Given a fiber oriented in direction θ after growth, Θ will depict the angle of this fiber before growth (see Fig. 3.1). Under the affine deformation assumption, the conservation of the cellulose mass that is initially contained within a control volume dV (dv after deformation) and infinitesimal angular sector $[\Theta, \Theta + d\Theta]$, is given by:

$$\rho(\theta) d\theta dv = R(\Theta) d\Theta dV \quad (3.2)$$

which amounts to:

$$\rho(\theta) = J^{-1} R(\Theta) \frac{d\Theta}{d\theta} \quad (3.3)$$

where $J \stackrel{\text{def}}{=} \frac{dv}{dV} = \det \mathbf{f}_g \geq 1$ measures the variation of volume. By detailing $\frac{d\Theta}{d\theta}$ (see Appendix 3.A), one obtains:

$$\rho(\theta) = \left(\frac{\lambda_1^2 + \lambda_2^2}{2} - \frac{\lambda_2^2 - \lambda_1^2}{2} \cos 2\hat{\theta} \right)^{-1} R(\Theta) \quad (3.4)$$

where λ_1 and λ_2 are the eigenvalues of \mathbf{f}_g (with $\lambda_1 \geq \lambda_2$); and angle $\hat{\theta}$ is the phase between θ and the principal angle of \mathbf{f}_g . The behavior predicted by Eq. (3.4) is further analyzed in Section 3.3.1.

For the sake of simplicity, we do not consider any growth-induced loading of the fibers, that would imply that the strain of each fiber would not be solely a function of its orientation, but also of its age and mechanical history. For example, two fibers deposited (unloaded) at two different times and in the same orientation could in principle be loaded in a different manner, depending on how much the wall has grown between the two fiber insertions. Here the strain energy associated with each fiber reflects the macroscopic elastic strain of the material, only. This is equivalent to considering that fiber strain energy relaxes at a longer time scale, in the same fashion as the macroscopic medium. Using this assumption, one can use Eq. (2.6) to relate the fiber density to stiffness.

3.2.2 Simulation

Triangular membrane finite elements equipped with \mathbb{P}_1 -Lagrangian shape functions are used to model the walls. To compute equilibrium, the equation of motion on nodes is dynamically integrated until the pressure forces \mathbf{f}_{ext} and reaction forces \mathbf{f}_{int} balance. Equilibrium is found when the maximal node force falls below a threshold. This step

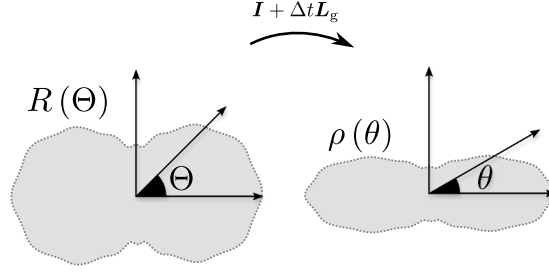


Figure 3.1: Deformation of an arbitrary angular distribution by an incremental deformation $\mathbf{f}_g = \mathbf{I} + \Delta t \mathbf{L}_g$.

is computed iteratively using the backward Euler method and conjugate gradient to solve intermediate linear systems. These are implemented in the open source software *Sofa* (Faure et al., 2012).

The reaction forces are evaluated from the expression of the total elastic energy of the system:

$$U = \sum_{\tau \in \mathcal{T}} \epsilon A^{(\tau)} \Psi^{(\tau)} \quad (3.5)$$

where \mathcal{T} is the set of triangle elements; ϵ depicts the wall thickness (assumed to be uniform); $A^{(\tau)}$ is the surface area of triangle τ in the stress-free configuration. The strain energy density $\Psi^{(\tau)}$ at triangle τ is defined as:

$$\Psi^{(\tau)} \stackrel{\text{def.}}{=} \Psi(\mathbf{E}^{(\tau)}) = \frac{1}{2} \mathbf{E}^{(\tau)} : \mathbb{C}_w^{(\tau)} : \mathbf{E}^{(\tau)}. \quad (3.6)$$

By derivation, one obtains the reaction force $\mathbf{f}_{\text{int}}^{(n)}$ at node n associated with node position $\mathbf{q}^{(n)}$:

$$\mathbf{f}_{\text{int}}^{(n)} = -\frac{\partial U}{\partial \mathbf{q}^{(n)}} = -\sum_{\tau \in \mathcal{T}} \epsilon A^{(\tau)} \frac{\partial \Psi^{(\tau)}}{\partial \mathbf{q}^{(n)}}. \quad (3.7)$$

The nodal external loads (pressure forces) are computed according to:

$$\mathbf{f}_{\text{ext}}^{(n)} = \frac{P}{3} \sum_{\tau \in \mathcal{T}_n} a^{(\tau)} \mathbf{n}^{(\tau)} \quad (3.8)$$

where $\mathbf{n}^{(\tau)}$ ($\|\mathbf{n}^{(\tau)}\| = 1$) and $a^{(\tau)}$ are respectively the outward normal and surface area of triangle τ (in the current configuration); \mathcal{T}_n is the set of the triangle elements that contain node n ; P is the excess of pressure relative to the outside.

After equilibrium is reached, we compute the growth step. Two computational solutions can be used for this operation. The first one consists of considering each finite element in its reference configuration, isolated from its neighbors. After growing each element separately, a compatible geometry can be computed by reconnecting the elements. This solution was used in Bassel et al. (2014) for example. Alternatively, we can

modify the elastic deformation gradient $\mathbf{F}_e^{(\tau)}$ in order to compensate for a change in the reference shape of the elements, which, from a computational standpoint, is a simpler operation. This solution is employed here.

The state of the system at mechanical balance and at time step t – noted $\mathbf{M}(t)$ – is defined through the list of node positions, and element deformation gradients:

$$\mathbf{M}(t) \stackrel{\text{def.}}{=} \left(\left\{ \mathbf{q}^{(n)}(t) \right\}_{n \in \mathcal{N}}, \left\{ \mathbf{F}_e^{(\tau)}(t) \right\}_{\tau \in \mathcal{T}} \right) \quad (3.9)$$

where \mathcal{N} depicts the set of nodes. Our general problem is to compute the evolution of the system, *i.e.* computing $\mathbf{M}(t + \Delta t)$ from $\mathbf{M}(t)$. For this, we re-express Eq. (3.1) according to $\mathbf{F}_e(t)$, using Eq. (1.6), as detailed hereafter. Algorithm 1 and Fig. 3.2(a) summarize the procedure employed to update the system.

Starting from a given equilibrium $\mathbf{M}(t)$, we compute the rate of plastic growth $\mathbf{L}_g^{(\tau)}(t)$ from $\mathbf{F}_e^{(\tau)}(t)$ for all triangles τ (using Eq. (1.12)), which allows to compute $\mathbf{F}_g^{(\tau)}(t + \Delta t)$ by Eq. (3.1). The total deformation gradient can now be expressed according to the new irreversible deformation $\mathbf{F}_g^{(\tau)}(t + \Delta t)$:

$$\mathbf{F}^{(\tau)}(t) = \mathbf{F}_e^{(\tau)}(t) \cdot \mathbf{F}_g^{(\tau)}(t) = \tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t) \cdot \mathbf{F}_g^{(\tau)}(t + \Delta t) \quad (3.10)$$

where $\tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t)$ is given by:

$$\tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t) \stackrel{\text{Eq. (3.1)}}{=} \mathbf{F}_e^{(\tau)}(t) \left(\mathbf{I} + \Delta t \mathbf{L}_g^{(\tau)}(t) \right)^{-1}. \quad (3.11)$$

Tensor $\tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t)$ defines the new elastic deformation of the system after application of growth. The tilde notation used in Eq. (3.11) emphasizes that, *a priori*, the state $\tilde{\mathbf{M}}(t + \Delta t) \stackrel{\text{def.}}{=} \left(\left\{ \mathbf{q}^{(n)}(t) \right\}_{n \in \mathcal{N}}, \left\{ \tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t) \right\}_{\tau \in \mathcal{T}} \right)$ is *not* at mechanical equilibrium. Instead, it is used as an initial condition to compute the next equilibrium $\mathbf{M}(t + \Delta t)$ (Algorithm 1).

After $\mathbf{L}_g^{(\tau)}(t)$ is computed for each triangle, the growth-induced fiber transport is applied provided the incremental deformation $\mathbf{f}_g^{(\tau)}(t)$ at step t , and by using Eq. (3.4). Since Eq. (3.4) is expressed in the angular domain (rather than the frequency domain) we first compute the *Inverse Fast Fourier Transform* of the CMF distribution $R^{(\tau)}$, then the *Fast Fourier Transform* to update the Fourier coefficients. In practice, the angular distributions $\rho^{(\tau)}$ and $\phi^{(\tau)}$ are approximated by their first 30 Fourier coefficients (Eqs. (2.22) and (2.33)). Although this operation is the costliest operation of the growth step, the computation of the mechanical equilibrium is by far the most important bottleneck of the full algorithm.

The last step consists of computing CMF deposition (according to Eq. (2.42)) and update the stiffness tensor (Eqs. (2.4) to (2.6)). This is done by the classical Runge-Kutta (RK4) procedure (assuming constant stress during the integration step).

For better spatial regularity of in the CMF density, the stress computed at each element is smoothed over its neighbors in a simple manner. The smoothed stress is used to evaluate Eq. (2.11). Since the PK2 stress is expressed in the 2D material coordinate system, it cannot be directly used to perform the smoothing. This is done for each triangle τ , by computing the stress $\sigma^{(\tau)}$ in the current configuration (Cauchy stress) from the PK2 stress $S^{(\tau)}$:

$$\sigma^{(\tau)} \stackrel{\text{def.}}{=} \frac{1}{\det \mathbf{F}_e^{(\tau)}} \mathbf{F}_e^{(\tau)} \cdot \mathbf{S}^{(\tau)} \cdot \left(\mathbf{F}_e^{(\tau)} \right)^T \quad (3.12)$$

Then, a mean stress is computed over the element neighborhood (weighted by triangle area). This stress, is generally of rank 3, as the neighborhood may be slightly non planar. It is then re-projected onto the mesh triangles and pulled back to the reference configuration, providing a smoothed PK2 stress field.

Algorithm 1 Growth increment

Require:

$M = (x, \mathbf{F}_e)$ at equilibrium

Ensure:

$M \leftarrow \text{Grow}(M)$ is the new equilibrium configuration after growth

- 1: $\mathbf{E} \leftarrow 1/2 (\mathbf{F}_e^T \cdot \mathbf{F}_e - \mathbf{I})$ \triangleright Elastic strain from the current equilibrium configuration
 - 2: $\mathbf{L}_g \leftarrow \Phi(\mathbf{E} - \mathbf{E}_y)_+$ \triangleright Strain-based growth
 - 3: $\mathbf{F}_e \leftarrow \mathbf{F}_e \cdot (\mathbf{I} + \Delta t \mathbf{L}_g)^{-1}$ \triangleright Redefining elastic deformation. Leaving equilibrium
 - 4: $M \leftarrow \text{Equilibrium}(x, \mathbf{F}_e)$ $\triangleright (FEM)$
-

3.2.3 Meshes

Surface meshes (for example those shown in Figs. 3.2(c) to 3.2(e) and 3.3(a)) were generated using the open source softwares *Blender* (blender.org), to sketch the initial shape, and *Graphite* (alice.loria.fr/software/graphite), that implements FEM-dedicated mesh enhancement routines.

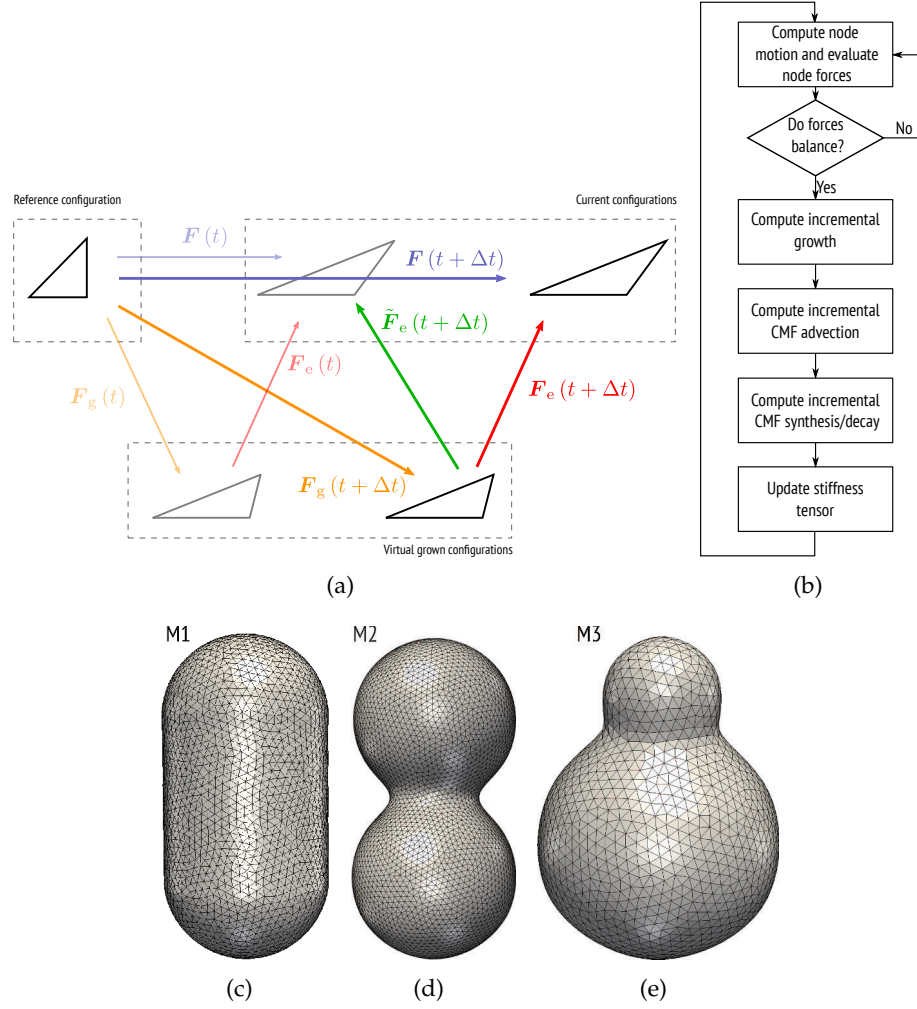


Figure 3.2: (a) Schematic representation of the growth procedure applied to a given triangle element. (b) General simulation pipeline. (c, d) Surface meshes: (c) Cylinder closed with hemispherical caps. (d, e) Two partially merged spheres.

3.3 Results

3.3.1 Verification of the model behavior

Convergence of the growth method In this section, we investigate the convergence of the growth method with respect to mesh resolution. In fact, if the solution excessively depends on the discretization, the reliability of the algorithm in accurately resolving the continuous model is not guaranteed. For this, we test the algorithm using spherical meshes with increasing resolution, from 40 to 5,000 nodes (Fig. 3.3(a)), in order to test to what extent an increase in mesh quality affects the various simulated properties. As we augment the mesh resolution, convergence is achieved as soon as the solution is no longer significantly affected by an improvement of the mesh quality.

For each version of the sphere, we simulated growth until the mesh surface area reached twice its initial value. For a given quantity p , we define (for each mesh m) the following measure of the deviation of p_m relative to a reference, at time t :

$$\varepsilon_p^m(t) = \frac{|p_m(t) - p_{\text{ref}}(t)|}{p_{\text{ref}}(t)} \quad (3.13)$$

where $p_{\text{ref}}(t)$ is the value associated with the reference mesh (the finest mesh of our set) at time t . We monitored the mesh radius² (ε_R), mean strain (ε_E), average CMF concentration (ε_{CMF}) and isotropy³ ($\varepsilon_{\text{isotropy}}$).

The results are given in Fig. 3.3(b). For extremely coarse meshes (e.g. the two first meshes on Fig. 3.3(a)), the error (for all the considered quantities) is confined to less than 10%. For finer meshes, the error falls below 1%, which we can consider as a satisfactory error for the present study. Figure 3.3(b), shows that the error tends to zero as a decreasing function of the mesh quality. This indicates that the growth algorithm (considered without stress feedback) in the case of a spherical shell, is convergent with respect to the mesh resolution. In the sequel, we employ meshes with an apparent resolution roughly comparable to that of the third mesh on Fig. 3.3(a).

²Computed as $R \stackrel{\text{def.}}{=} 3 \frac{\text{Volume}}{\text{Area}}$.

³With isotropy $\stackrel{\text{def.}}{=} 1 - \langle |\hat{\rho}_1| / \hat{\rho}_0 \rangle_{\text{triangles}}$, see Eq. (2.22).

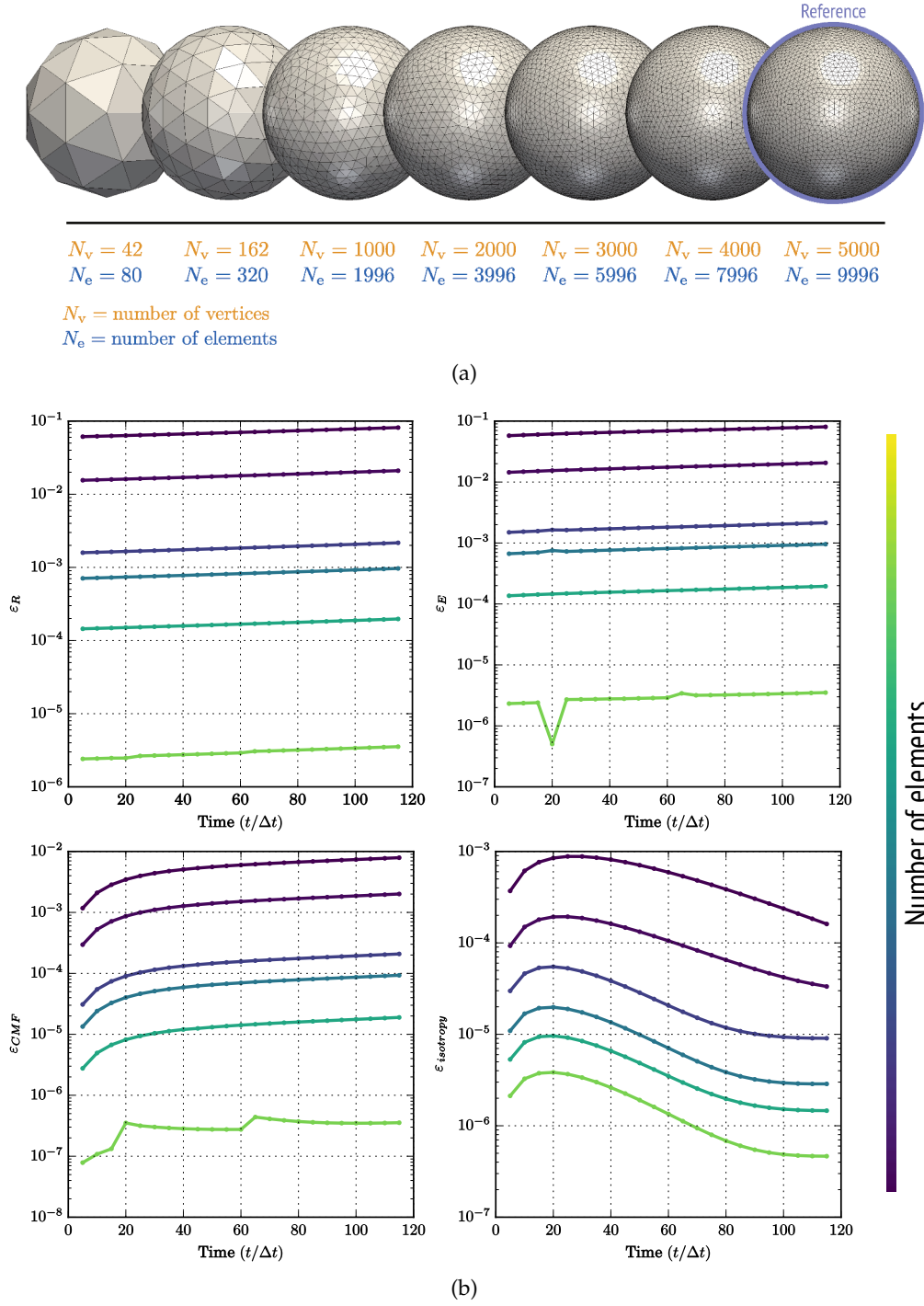


Figure 3.3: **Convergence of the growth method** (a) Various meshes of a sphere with increasing resolution. (b) Error in radius (ϵ_R), strain (ϵ_E), CMF concentration (ϵ_{CMF}) and isotropy $\epsilon_{isotropy}$ presented in the main text (Eq. (3.13)). Simulation parameters: turgor pressure $P = 0.1$ MPa, mesh typical length = $40 \mu\text{m}^3$, wall extensibility $\Phi = 1 \text{ s}^{-1}$, growth threshold $E_y = 2\%$, $\gamma = 0$, wall thickness $\epsilon = 0.4 \mu\text{m}$, cellulose deposition constant $k_\rho = 1 \text{ mol} \cdot \mu\text{m}^{-3} \cdot \text{s}^{-1}$, cellulose decay constant $k'_\rho = 1 \text{ s}^{-1}$, surface fiber Young's modulus $\epsilon Y_f = 40 \text{ MPa} \cdot \mu\text{m}$ and reduced surface Young's modulus of the gel $\epsilon Y \approx 1 \text{ MPa} \cdot \mu\text{m}$, matrix Poisson's ratio $\nu = 0.2$.

Fiber advection We see from Eq. (3.4), that the CMFs tend to accumulate in the direction of main deformation, and are depleted in the perpendicular direction. This captures a tendency of CMFs to passively align with the direction of main growth. In addition, the model captures a dilution of the CMFs: these tend to locally vanish due to the increase in wall volume (see also Fig. 3.5). This can be evidenced from Eq. (3.4) in the particular case where R is an isotropic distribution (say equal to R^0). In this case, one may verify that the dilution factor is the variation of volume J :

$$\int_{\pi} \rho = R^0 \int_{\pi} \left(\frac{\lambda_1^2 + \lambda_2^2}{2} - \frac{\lambda_1^2 - \lambda_2^2}{2} \cos 2\hat{\theta} \right)^{-1} d\theta = \frac{\pi R^0}{J} \quad (3.14)$$

(see Appendix 3.B).

Figure 3.4 shows minimal examples of these behaviors, considering a given distribution subject to an arbitrary anisotropic and steady growth field. In a more general case, *i.e.* where growth depends itself on fiber distributions (Fig. 3.5), we use FEM simulations. Figure 3.5 shows simulations of growing pressurized membranes (Figs. 3.2(d) and 3.2(d)) with initial uniform and isotropic CMF distribution, where stress feedback, as well as cellulose turnover are inactivated (e.g. $k_{\rho} = k'_{\rho} = \gamma = 0$). Under this assumption, we verify that the total mass of cellulose is a conserved quantity (as shown by the dotted line in Fig. 3.5(b)). Moreover, we observe that the decrease of cellulose concentration is more rapid in fast growing zones (Figs. 3.5(c) and 3.5(d)), and that the CMFs tend to passively orient in the main direction of growth in the concave part of the mesh, where growth is anisotropic (Figs. 3.5(c) and 3.5(d)).

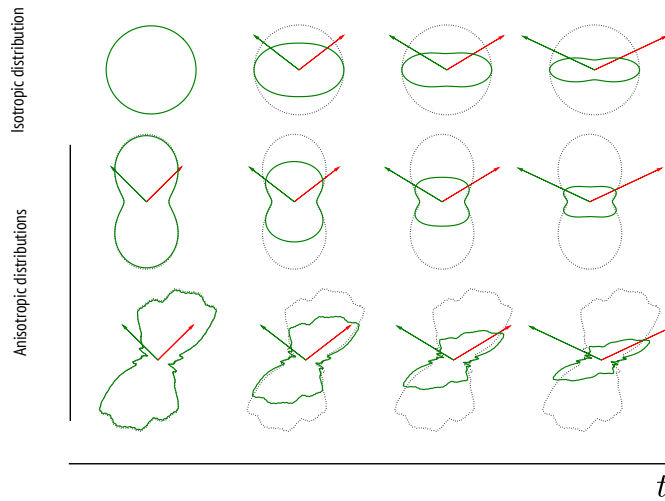


Figure 3.4: Advection of a fiber distribution by a steady unidirectional growth. The arrows indicate the successive deformations of an initial orthonormal basis. Plain and dashed lines show the current and initial polar plots of the distribution. From top to bottom: initially isotropic distribution, anisotropic distribution orthogonal to the growth axis, general distribution.

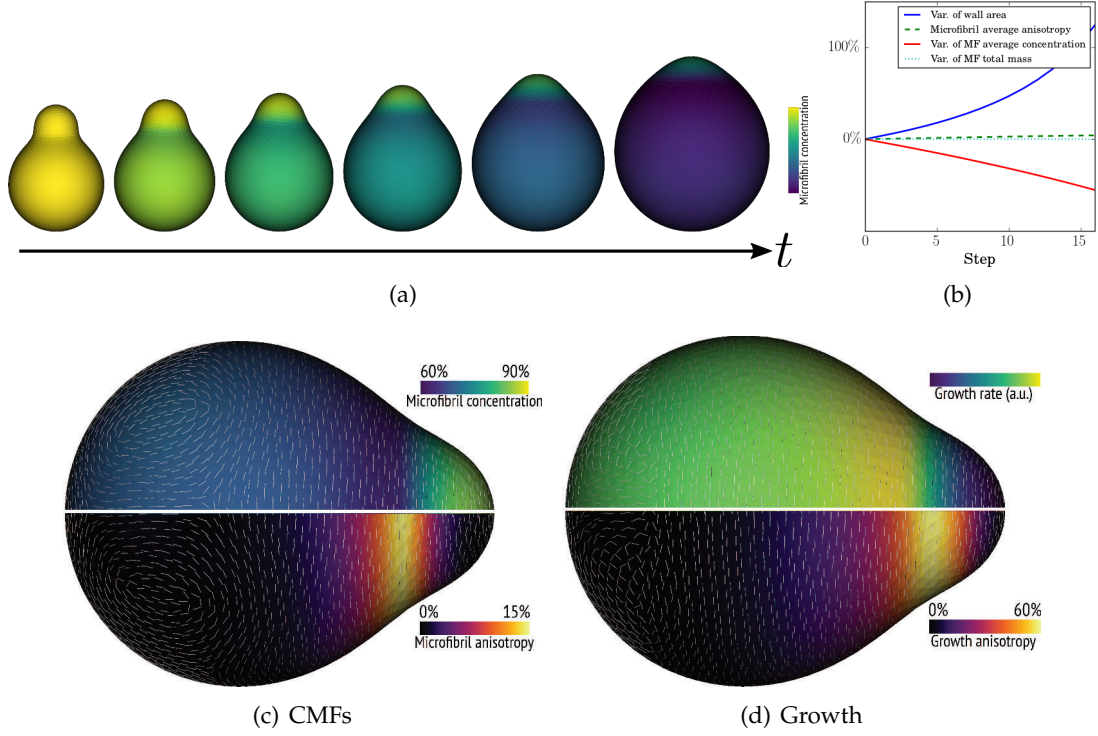


Figure 3.5: **Coupling growth and CMF transport in a FEM simulation of growth.** Growth simulation for mesh M3 (Fig. 3.2(e)). The simulation is initialized with uniform isotropic CMF density. No CMF polymerization/depolymerization are considered ($k_p = k'_p = 0$). (a) CMF concentration at various time steps. (b) Variation ($\text{value}(t)/\text{value}(0) - 1$) of wall area, of average CMF concentration, of CMF total mass and of CMF average anisotropy. (c, d) Simulation result after several growth steps. (c) CMF density (as a % of the initial density) and anisotropy. Rods indicate direction of main alignment. (d) Growth rate and anisotropy. Rods indicate main growth axis. This simulation confirms the depletion of cellulose in fast growing zones, and the tendency of CMFs to rotate in the direction of main growth. Simulation parameters: turgor pressure $P = 0.1$ MPa, mesh typical length = $100 \mu\text{m}$, wall extensibility $\Phi = 1 \text{ s}^{-1}$, growth threshold $E_y = 1\%$, $\gamma = 0$, reduced surface fiber Young's modulus $\epsilon Y_f = 100 \text{ MPa} \cdot \mu\text{m}$ and reduced surface Young's modulus of the gel $\epsilon Y \approx 1 \text{ MPa} \cdot \mu\text{m}$, matrix Poisson's ratio $\nu = 0.2$.

3.3.2 Growth of stress-adaptive shells

General idea In Chapter 2, we suggested that the stress feedback may be a mechanism to amplify shape asymmetry. For this reason, it may have the potential to promote symmetry breaking, by amplifying small perturbations in the initial shape. To test this behavior, we simulated the growth of simple abstract surfaces, with various degrees of stress feedback. We considered key parameters of the stress feedback, that are the sensitivity coefficient γ (that governs the sharpness of CMT alignment, see Eq. (2.35)) and the kinetic coefficients k_p and k'_p that govern cellulose turnover (for the sake of

simplicity we impose $K_\rho = k_\rho/k'_\rho = 1 \text{ mol} \cdot \mu\text{m}^{-3}$, which allows to vary only one parameter). Parameter k'_ρ governs the rapidity with which the tissue modifies its stiffness in response to a stress, and characterizes the typical time of CMF reorganization. Intuitively, a sensitive CMT system coupled with a very slow CMF dynamics will fail in rapidly responding to stress by modifying stiffness. Conversely, we may expect that a rapid cellulose renewal will be of little interest if CMTs insufficiently coalign along stress main axis.

As in Chapter 2, we considered pressurized membrane meshes (no inner tissues). Initially, the cellulose system is set in the trivial null-stress steady concentration, which mostly depends on K_ρ (see Eq. (2.43)). Hence, all simulations are initialized with the same isotropic elasticity. Simulations are conducted until the mesh surface area reaches twice its initial value.

Spherical mesh The most symmetric shape is the sphere. Evidently, by a trivial symmetry argument, the sphere is a fixed point of the system. We here study its stability.

We simulated stress feedback and growth using a spherical mesh composed of about 4,000 nodes and 8,000 triangular elements. The results of the simulations on the sphere are shown in Fig. 3.6. Confirming the intuition given above, we verify that for a slow and/or insensitive feedback, growth is uniform and spherical, which means that the sphere is stable for this range of parameters. By contrast, by using a fast and sensitive feedback (upper-right corner of Fig. 3.6), we observed strong growth heterogeneities due to rapid anisotropization of the tissue, suggesting that the spherical shape will be unstable for this range of parameters. Yet, in absence of a remeshing procedure, this kind of heterogeneities occurred with a rapid deterioration of the mesh quality, as quantified by the histograms in Fig. 3.6, showing the distributions of triangle skewness and mesh roughness (these quantities are defined in Appendix 3.C). For extremely fast and sensitive feedback, maximum skewness and roughness in fact exceeded 50%.

To assess the nature of this apparent “patterning” in the growth rate, we tested the sensitivity of this result with respect to mesh quality, using various spherical meshes with a node number ranging between 3,000 and 20,000 (see Fig. 3.7). At the end of the simulation, we quantified for each mesh the *discrete correlation length* (defined in Appendix 3.D) which provides an estimate of the typical length of the heterogeneity. For all simulations, this length was comparable to the typical size of the triangular elements, although the number of elements varied significantly between all meshes (see Fig. 3.7). This means that, for this range of extreme (potentially non-physiological) parameters, our FEM pipeline fails in converging with respect to mesh size. Note also

that the smoothing operation presented in Section 3.2.2 is performed over the immediate neighbors of a given triangle. Therefore the smoothing radius also varied with mesh size. In future work, we should investigate a more sophisticated, potentially mesh-independent stress-smoothing procedure. Procedures to improve the quality of the computed stress should be also investigated.

Quasi-spherical spheroids The previous example suggests that the spherical shape is globally stable, although there exists a range of parameters of the feedback that will probably lead to instabilities (confirming this result would require further numerical investigations). The only perturbations considered in that case were those introduced by the meshing, which approximates a sphere by a polyhedron. To study the effect of larger perturbations, we considered the case of almost-spherical spheroids: a *prolate* spheroid (elongated sphere) and *oblate* spheroid (flattened sphere). In both cases, we considered a 5% perturbation in diameter.

The stress field in both cases was computed (see Fig. 3.8), showing that in the prolate case, hoop stress slightly dominates, while in the oblate case, the dominating stress is mostly axial. This can be explained by the different natures of the curvature in these two shapes, although both are very close to a sphere. A stress feedback has then the potential to yield utterly different behaviors depending on whether the perturbation is prolate or oblate.

In the case of the slightly prolate spheroid (Fig. 3.9), the system remained roughly spherical provided a weak feedback, as the material stiffness remained more or less isotropic. In the case of an extremely sharp and rapid feedback, growth became highly heterogeneous and noisy as previously observed. A slightly weaker feedback (e.g. $\gamma = 10 \text{ MPa}^{-1}$, $k_p = 1 \text{ mol} \cdot \mu\text{m}^{-3} \cdot \text{s}^{-1}$) allowed for a response to the slight initial stress anisotropy (as CMTs aligned circumferentially) and for the amplification of the initial shape asymmetry. In fact, the shape elongated as a cylinder. Here the mesh skewness increased while, comparatively, mesh roughness was small, which is the mark of a cohesive anisotropic growth.

In the case of the slightly oblate spheroid (Fig. 3.10) the system again developed towards a sphere in the case of a slow or insensitive feedback. Spheroid flatness was slightly amplified for more sensitive and faster feedback, yet this often occurred together with a slight elongation of the shape in the axial plane, that is the plane orthogonal to the initial symmetry axis (not shown). This suggests that the feedback tends to destabilize flat shapes by breaking the initial axial symmetry, and promotes elongation. In these simulations, the stress feedback tended to elongate triangles in a very

heterogeneous manner (as shown by the histograms in Fig. 3.10). In particular, singular growth spots appeared at the poles of the structure.

Growth of highly asymmetric shapes In the previous paragraph, we saw that a slightly prolate sphere may develop axially. We next confirmed this result on a more elongated shape (a cylindrical "pill", shown in Fig. 3.2(c)). In this case, axial cylindrical expansion was promoted by a sharp feedback ($\gamma > 15 \text{ MPa}^{-1}$), see Fig. 3.11(a), as we observed an exclusively-axial growth, resulting in a cylindrical expansion with quasi-constant radius. By contrast, lower feedback sensitivity resulted in a radial expansion.

A similar set of simulations, starting from a shape displaying a concave zone (here the shape shown in Fig. 3.2(d)) also confirmed the suggestion made in Chapter 2, that a stress feedback is a potential way to maintain a concave frontier (see Fig. 3.11(b)). In fact, for sufficient values of γ , we observed that the frontier radius barely increased, while the rest of the tissue dilated substantially. This may be a way to amplify the growth heterogeneities between two developmental zones.

Simulation of highly oblate shapes (flat spheroids) generally did not properly converge, even for extremely fine meshes and extremely small time step size. This may partly be due to the emergence of compressive stresses and buckling at the ridge of the spheroid. Membrane finite elements (that do not take bending energy into consideration) are not appropriate to represent this kind of effect. We further discuss this special case – in particular its biological relevance – in the discussion of this chapter.

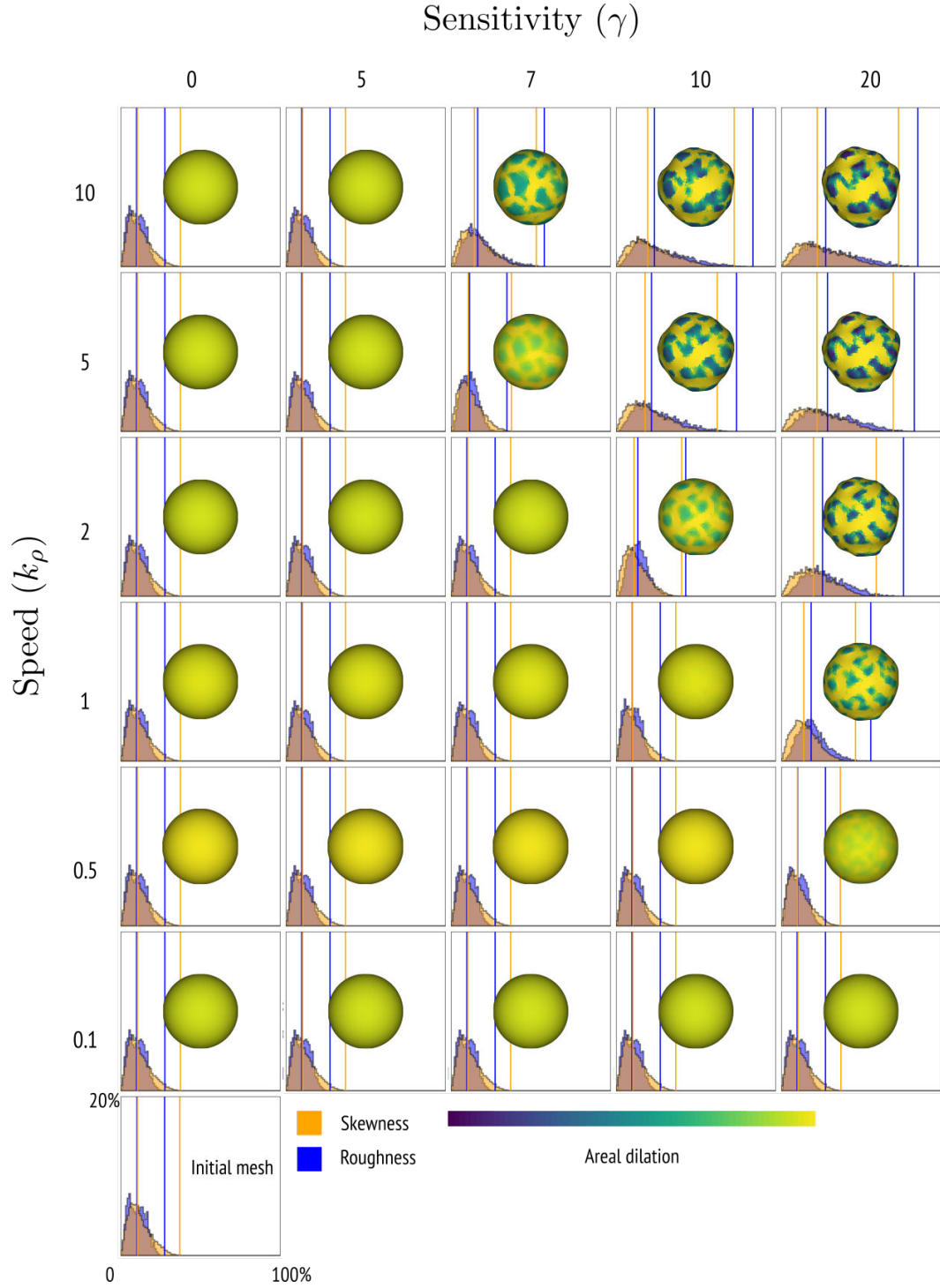


Figure 3.6: Growth of a stress-adaptive sphere. Growth is computed until surface area is twice the initial one, for various values of the sensitivity coefficient γ and kinetic constant k_ρ (taking $K_\rho = 1 \text{ mol} \cdot \mu\text{m}^{-3}$). Heatmap shows the areal dilation of triangles (namely the ratio between final area vs initial area). Histograms quantify mesh quality at the end of the simulation: skewness and roughness (Appendix 3.C) are respectively represented in orange and blue (vertical lines show median and maximum values). Simulation parameters: turgor pressure $P = 0.1 \text{ MPa}$, mesh typical length = $40 \mu\text{m}$, wall extensibility $\Phi = 1 \text{ s}^{-1}$, growth threshold $E_y = 1\%$, $\gamma = 0$, wall thickness $\epsilon = 0.4 \mu\text{m}$, reduced surface fiber Young's modulus $\epsilon Y_f = 40 \text{ MPa} \cdot \mu\text{m}$ and reduced surface Young's modulus of the gel $\epsilon Y \approx 1 \text{ MPa} \cdot \mu\text{m}$, matrix Poisson's ratio $\nu = 0.2$.

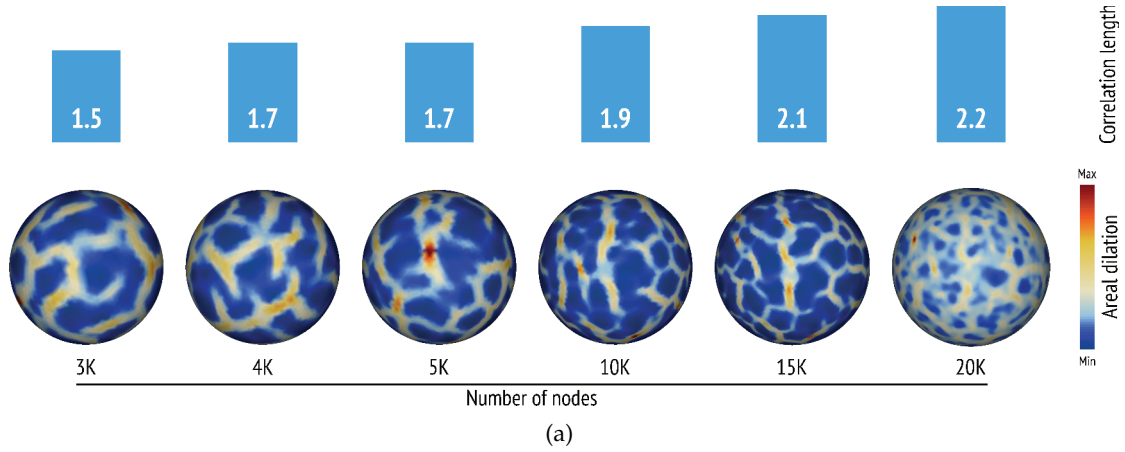


Figure 3.7: Growing sphere equipped with fast and sharp feedback: $\gamma = 20 \text{ MPa}^{-1}$ and $k_\rho = 5 \text{ mol} \cdot \mu\text{m}^{-3}$, $k'_\rho = 5$ (all other parameters are equal to those used in Fig. 3.6). Areal dilation (triangle current area divided by initial area, interpolated at nodes) shown in the initial configuration and discrete correlation length (see Appendix 3.D) for increasing mesh resolution.

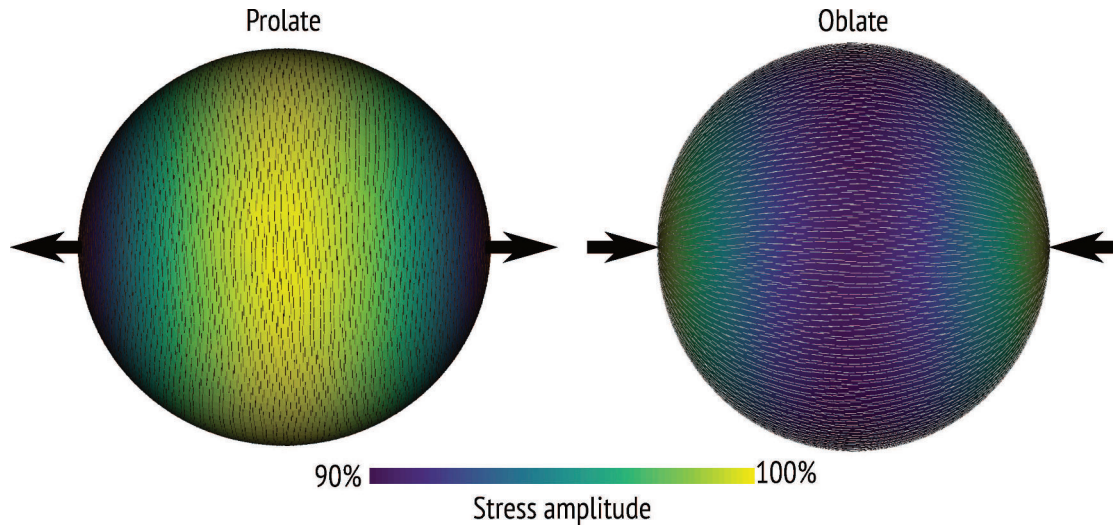


Figure 3.8: Stress at equilibrium in the oblate and prolate spheroids. Rods and colormap respectively indicate the main axis and trace of the Cauchy stress. Arrows indicate the axis of the shape elongation/flattenning.

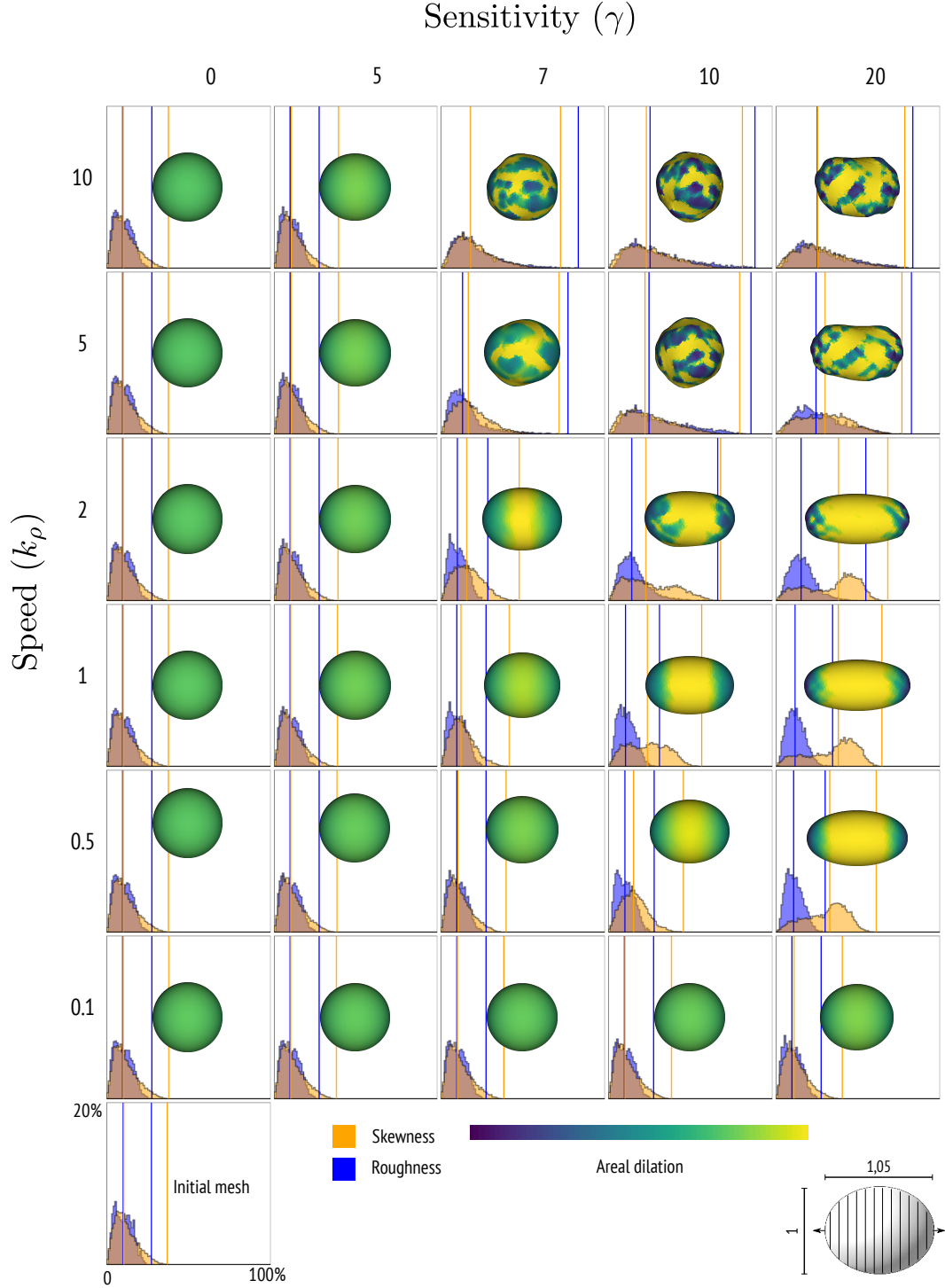


Figure 3.9: 5% prolate spheroid. Growth is computed until surface area is twice the initial one, for various values of the sensitivity coefficient γ and kinetic constant k_ρ (taking $K_\rho = 1 \text{ mol} \cdot \mu\text{m}^{-3}$). Heatmap shows the areal dilation of triangles (namely the ratio between final area vs initial area). Histograms quantify mesh quality at the end of the simulation: skewness and roughness (Appendix 3.C) are respectively represented in orange and blue (vertical lines show median and maximum values). Simulation parameters: turgor pressure $PP = 0.1 \text{ MPa}$, mesh typical length = $40 \mu\text{m}^3$, wall extensibility $\Phi = 1 \text{ s}^{-1}$, growth threshold $E_\gamma = 3\%$, surface fiber Young's modulus $\epsilon Y_f = 40 \text{ MPa} \cdot \mu\text{m}$ and matrix Young's modulus $\epsilon Y \approx 1 \text{ MPa} \cdot \mu\text{m}$, matrix Poisson's ratio $\nu = 0.2$.

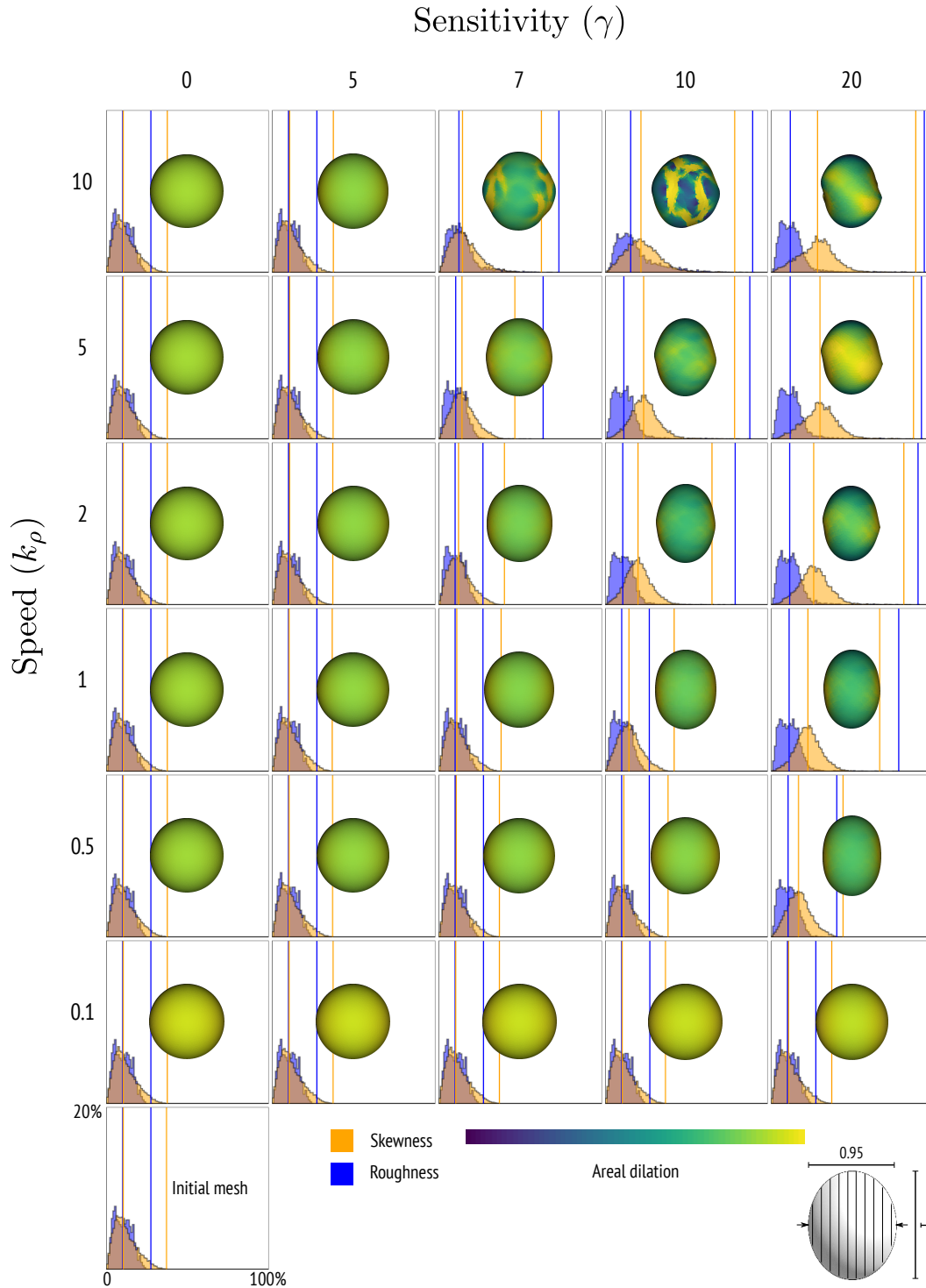


Figure 3.10: 5% oblate spheroid. Growth is computed until surface area is twice the initial one, for various values of the sensitivity coefficient γ and kinetic constant k_ρ (taking $K_\rho = 1 \text{ mol} \cdot \mu\text{m}^{-3}$). Heatmap shows the areal dilation of triangles (namely the ratio between final area vs initial area). Histograms quantify mesh quality at the end of the simulation: skewness and roughness (Appendix 3.C) are respectively represented in orange and blue (vertical lines show median and maximum values). Simulation parameters: turgor pressure $P = 0.1 \text{ MPa}$, mesh typical length $= 40 \mu\text{m}$, wall extensibility $\Phi = 1 \text{ s}^{-1}$, growth threshold $E_\gamma = 1\%$, wall thickness $\epsilon = 0.4 \mu\text{m}$, surface fiber Young's modulus $\epsilon Y_f = 40 \text{ MPa} \cdot \mu\text{m}$ and matrix Young's modulus $\epsilon Y \approx 1 \text{ MPa} \cdot \mu\text{m}$, matrix Poisson's ratio $\nu = 0.2$.

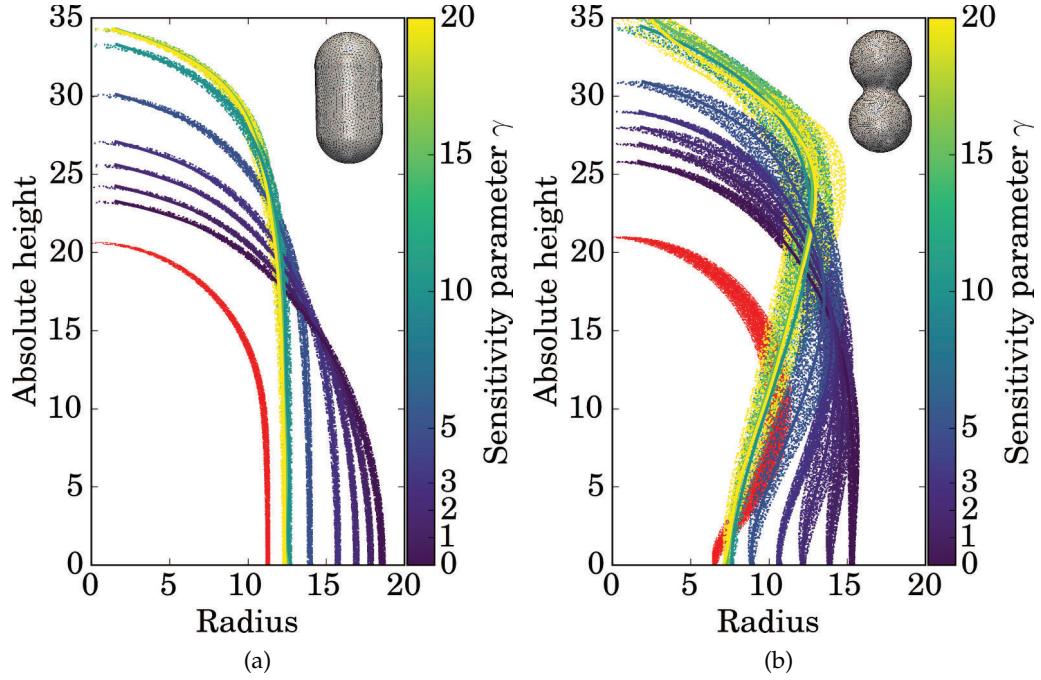


Figure 3.11: Growth profiles of (a) the closed cylinder (Fig. 3.2(c)) and (b) merged spheres (Fig. 3.2(d)), for various values of γ . Each dot represents the distance from symmetry axis and height of the mesh nodes. Bold lines represent a polynomial fit. Red dots show the profile taken after the first equilibrium is reached. The other profiles correspond to the configuration when the mesh surface area has reached twice its original surface area. Simulation parameters: turgor pressure $P = 0.1$ MPa, mesh typical length = $40 \mu\text{m}$, wall extensibility $\Phi = 1 \text{ s}^{-1}$, growth threshold $E_y = 2\%$, cellulose deposition constant $k_\rho = 1 \text{ mol} \cdot \mu\text{m}^{-3} \cdot \text{s}^{-1}$, cellulose decay constant $k'_\rho = 1 \text{ s}^{-1}$, reduced surface fiber Young's modulus $\epsilon Y_f = 40 \text{ MPa} \cdot \mu\text{m}$ and reduced surface Young's modulus of the gel $\epsilon Y \approx 1 \text{ MPa} \cdot \mu\text{m}$, matrix Poisson's ratio $\nu = 0.2$.

3.4 Discussion

Amplification of tissue asymmetry by a stress feedback In this chapter, we have studied to what extent a stress feedback can amplify initial perturbations in shape. This is possibly required in the establishment of highly asymmetric organs such as stems or leaves. *In vivo*, these organs start as small, more or less roundish bumps, which result from a local modification of the epidermal cell wall constitution (see [Sassi et al., 2014](#)). These bumps may further develop in a highly anisotropic manner.

To study how this transition in growth mode occurs, we used simple abstract shapes, *i.e.* perturbed spheres, as initial shapes. In particular, we employed the classic pressurized vessel assumption, that consists of neglecting the mechanical role of inner tissues and of considering only the outermost wall as an elastic membrane.

This theoretical study suggested that the stress feedback has the potential to promote axial elongation starting from a slightly prolate spheroid (Fig. 3.9), provided a reasonably rapid and sensitive feedback is active, and, secondly, to maintain axial growth while keeping a nearly constant radius (Fig. 3.11(a)). This result is consistent with what was found in [Bozorg \(2016\)](#), [Hamant et al. \(2008\)](#), that are based on more *ad hoc* stress feedback models. This amplification was less clear in the case of an oblate spheroid and flat expansion, the simulation of which was more difficult to conduct, due for example to fast and localized mesh deterioration (Fig. 3.10). Further numerical investigations – including the development of a proper remeshing algorithm – will be needed to validate this result. To date, the implementation of a triangle subdivision method is ongoing.

One can still expect, based on mere intuition, that a hollow membrane model will fail in producing very thin shapes. In principle, broadening should result in an increase of the stress in the flattest parts of the structure (namely at the poles), the curvature of which should tend to zero. In the most extreme case, the maintenance of an almost null curvature will require polar tissue tension to approach infinity, in order to balance turgor forces. This is of course very unlikely to happen in plants. In reality, part of this stress should be transferred to inner tissues, that are neglected in the pressure-vessel hypothesis. In such a scenario, one can expect that inner walls may have a crucial mechanical role in maintaining and amplifying flatness. Whether a stress feedback can account for this kind of shape is treated in the next chapter, in the case of *Arabidopsis thaliana* and tomato leaves and sepals that (as in many other species) are extremely asymmetric organs.

On the maintenance of a homeostatic cellulose density To achieve efficient shape amplification during the transient organ initiation, fast and sharp feedback may be necessary, as abstracted through the idealized case of a perturbed sphere. In our model, these features are represented through parameters γ and k'_ρ . We hereafter discuss the role and interpretation of k'_ρ from mathematical and biological standpoints.

The equation governing the dynamics of cellulose density (Eq. (2.7)) is composed of two terms in its right-hand side: a term representing the CMT-guided polymerization of cellulose, which models a rather well identified biological process (*i.e.* the CMT-guided deposition of cellulose by CSCs, see [Paredes et al., 2006](#)), and an isotropic linear decay. The latter was necessary in Chapter 2 to guarantee the existence of a steady cellulose concentration. However, its mathematical necessity is here less obvious, in so far as the growth-induced advection also contributes to reducing CMF concentration $c_\rho = \int_\pi \rho$. This can be expressed in the form of a classic advection-reaction equation (that directly derives from the continuity equation):

$$d_t c_\rho = k_\rho - \underbrace{k'_\rho c_\rho}_{\text{Cellulose decay}} - \underbrace{c_\rho \nabla \cdot \mathbf{v}}_{\text{Advection}} \quad (3.15)$$

(where d_t depicts the *material* derivative, see [Holzapfel, 2000](#)). This shows that growth has the potential to induce a decrease of cellulose concentration, yet in a passive manner (since \mathbf{v} is an emergent property of our system, in contrast to k'_ρ).

While previously, the steady regime depended upon the independent values of k_ρ and k'_ρ , advection depends upon the growth rate, and therefore is regulated through stiffness and CMF density itself, probably in a less direct manner than a kinetic coefficient like k'_ρ . One immediate question is then: can advection *only* (*i.e.* taking $k'_\rho = 0$) overcome cellulose deposition, so that the cellulose concentration can be maintained in a homeostatic, steady value?

Intuitively, the answer should be negative. Indeed, we may expect that such a system (advection + polymerization with constant rate) would be in principle hard to stabilize. A large value of k_ρ would most plausibly result in growth arrest, since stiffness would increase and, likewise, advection should vanish. Conversely, a low value of k_ρ would imply soft walls and fast growth, and therefore fast depletion of the fibers, potentially at the expense of tissue integrity. In addition, it is not certain that advection would allow a rapid turnover of the fibers allowing for a rapid response of the system to a change in stress.

To rule out this issue, we can picture two different parsimonious mechanisms. First,

we may keep the depolymerization term $k'_\rho \rho(\theta)$ used in Eq. (2.7), which allows an arbitrarily rapid saturation of cellulose concentration and provides flexibility to the system. This was the solution chosen in this chapter and the previous one. This term can reflect a depolymerization process, possibly controlled by the cell through some enzymatic activity. To date however, no such depolymerization mechanism has yet been identified to our knowledge. In the alternative scenario, cellulose polymerization kinetics could depend itself upon growth rate and/or current cellulose concentration. In other terms, the cell may dynamically regulate the quantity of cellulose it synthesizes by probing its growth and/or structural state and controlling k_ρ accordingly. This might also be a plausible scenario, supported by the recent identification of several potential wall integrity and growth sensors like THESEUS or FERONIA involved in the control of cellulose secretion (Cheung and Wu, 2011, Hématy and Höfte, 2008, Verger and Hamant, 2018, Wolf et al., 2014). However, in this scenario, it is not clear how a cell could rapidly renew its wall composition and efficiently perform a change in its growth behavior. Possibly, both scenarios may be actually valid.

Note that we assumed that growth rate depends on fiber concentration mostly. This is a coarse approximation, as, for example, growth is also mediated through the hemi-cellulose tethers that connect the fibers together (Cosgrove, 2005, Geitmann and Ortega, 2009) as modeled in Dyson et al. (2012). These were here abstracted through the isotropic elasticity tensor \mathbb{C}_g . To fully understand the loop between wall remodeling and growth, and its integration in global morphogenesis, finer models of the walls and its growth, compatible with a global simulation of mechanics, are required.

Appendices

3.A Proof of Eq. (3.4)

I here detail the establishment of Eq. (3.4) that describe the deformation of the CMF distribution due to the deformation increment $\mathbf{f}_g = \mathbf{I} + \Delta t \mathbf{L}_g^n$.

In particular we determine an expression for $\frac{d\Theta}{d\theta}$ involved in the mass conservation equation (Eq. (2.24)). By definition:

$$\Theta = \arctan 2 (\mathbf{j} \cdot \mathbf{N}, \mathbf{i} \cdot \mathbf{N}) = \arctan 2 (\mathbf{j}^* \cdot \mathbf{n}, \mathbf{i}^* \cdot \mathbf{n}) \quad (3.16)$$

where $\mathbf{i}^* = \mathbf{f}_g^{-T} \cdot \mathbf{i}$ and $\mathbf{j}^* = \mathbf{f}_g^{-T} \cdot \mathbf{j}$. Recalling that $\nabla \arctan 2 (y, x) = \left(\frac{-y}{x^2+y^2}, \frac{x}{x^2+y^2} \right)$, one obtains, by differentiating Eq. (3.16):

$$\begin{aligned} \frac{d\Theta}{d\theta} &= \frac{d\mathbf{n}}{d\theta} \cdot \frac{d\Theta}{d\mathbf{n}} \\ &= \mathbf{n}_{\frac{\pi}{2}} \cdot \frac{-(\mathbf{j}^* \cdot \mathbf{n}(\theta)) \mathbf{i}^* + (\mathbf{i}^* \cdot \mathbf{n}(\theta)) \mathbf{j}^*}{\mathbf{i}^* \cdot \mathbf{n}(\theta)^2 + \mathbf{j}^* \cdot \mathbf{n}(\theta)^2} \quad \text{with} \quad \mathbf{n}_{\frac{\pi}{2}} = \mathbf{n}(\theta + \pi/2) \\ &= \mathbf{n}_{\frac{\pi}{2}} \cdot \left(\frac{(\mathbf{i}^* \otimes \mathbf{j}^*)^T - \mathbf{i}^* \otimes \mathbf{j}^*}{\|\mathbf{f}_g^{-1} \cdot \mathbf{n}\|^2} \cdot \mathbf{n} \right) \\ &= \frac{\mathbf{i}^* \otimes \mathbf{j}^* - (\mathbf{i}^* \otimes \mathbf{j}^*)^T}{\|\mathbf{f}_g^{-1} \cdot \mathbf{n}\|^2} \cdot \mathbf{n}_{\frac{\pi}{2}} \cdot \mathbf{n} \\ &= \frac{\mathbf{i}^* \otimes \mathbf{j}^* - (\mathbf{i}^* \otimes \mathbf{j}^*)^T}{\|\mathbf{f}_g^{-1} \cdot \mathbf{n}\|^2} : \mathbf{n} \otimes \mathbf{n}_{\frac{\pi}{2}}. \end{aligned}$$

From mass conservation (Eq. (2.24)), one derives:

$$\rho(\theta) = R(\Theta) \frac{\lambda(\theta)^2}{J} \left(\mathbf{i}^* \otimes \mathbf{j}^* - (\mathbf{i}^* \otimes \mathbf{j}^*)^T \right) : \mathbf{n} \otimes \mathbf{n}_{\frac{\pi}{2}}. \quad (3.17)$$

where Θ is obtained from Eq. (3.16) and where $\lambda(\theta) = 1/\|\mathbf{f}_g^{-1} \mathbf{n}\|$ is the stretch in direction θ .

We can assume, without loss of generality that \mathbf{f}_g is diagonal in \mathcal{B} (\mathbf{f}_g is a symmetric tensor):

$$[\mathbf{f}_g]_{\mathcal{B}} = \begin{bmatrix} \lambda_1 & 0 \\ 0 & \lambda_2 \end{bmatrix} \quad (3.18)$$

Hence:

$$\mathbf{i}^* \otimes \mathbf{j}^* = (\mathbf{f}_g^{-1} \cdot \mathbf{i}) \otimes (\mathbf{f}_g^{-1} \cdot \mathbf{j}) = \frac{1}{J} \mathbf{i} \otimes \mathbf{j} \quad (3.19)$$

giving:

$$(\mathbf{i}^* \otimes \mathbf{j}^* - (\mathbf{i}^* \otimes \mathbf{j}^*)^T) : \mathbf{n} \otimes \mathbf{n}_{\frac{\pi}{2}} = \frac{1}{J}. \quad (3.20)$$

In addition:

$$\lambda(\theta)^2 = \frac{1}{\mathbf{f}_g^{-2} : \mathbf{n} \otimes \mathbf{n}} = \frac{1}{\left(\frac{\cos \theta}{\lambda_1}\right)^2 + \left(\frac{\sin \theta}{\lambda_2}\right)^2} = \frac{\frac{2\lambda_1^2\lambda_2^2}{\lambda_1^2 + \lambda_2^2}}{1 - \frac{\lambda_1^2 - \lambda_2^2}{\lambda_1^2 + \lambda_2^2} \cos(2\theta)}. \quad (3.21)$$

Combining the two previous expressions with Eq. (3.17) finally gives:

$$\rho(\theta) = \left(\frac{\lambda(\theta)}{J}\right)^2 R(\Theta) = \frac{\frac{2}{\lambda_1^2 + \lambda_2^2}}{1 - \frac{\lambda_1^2 - \lambda_2^2}{\lambda_1^2 + \lambda_2^2} \cos(2\theta)} R(\Theta). \quad (3.22)$$

For the previous expression to be valid in any basis, the argument of the cosine must take into account the phase shift that corresponds to the angle $\theta_{\mathbf{f}_g}$ of main deformation. This is done by replacing θ by $\hat{\theta} = \theta - \theta_g$ in the previous equation, where θ_g is the angle of main growth.

3.B Proof of Eq. (3.14)

Let's note $A = \frac{\lambda_1^2 + \lambda_2^2}{2}$ and $\alpha = \frac{\lambda_1^2 - \lambda_2^2}{\lambda_1^2 + \lambda_2^2}$ within the scope of this appendix. We compute:

$$\begin{aligned}
 \int_{\pi} \frac{d\theta}{A(1 - \alpha \cos 2\theta)} &= \frac{1}{2A} \int_{2\pi} \frac{d\theta}{1 - \alpha \cos 2\theta} \\
 &= \frac{1}{2A} \int_{2\pi} \frac{d\theta}{1 + \alpha - 2\alpha \cos^2 \theta} \\
 &= \frac{2}{A} \int_0^{\frac{\pi}{2}} \frac{\sec^2 \theta d\theta}{(1 + \alpha) \sec^2 \theta - 2\alpha} \quad \left(\sec \theta = \frac{1}{\cos \theta} \right) \\
 &= \frac{2}{A(1 + \alpha)} \int_0^{\frac{\pi}{2}} \frac{\sec^2 \theta d\theta}{\frac{1 - \alpha}{1 + \alpha} + \tan^2 \theta} \quad (\sec^2 \theta = 1 + \tan^2 \theta) \\
 &= \frac{2}{A(1 + \alpha)} \int_0^{\infty} \frac{dx}{\frac{1 - \alpha}{1 + \alpha} + x^2} \quad (\text{subst.: } x = \tan \theta) \\
 &= \frac{\pi}{A\sqrt{1 - \alpha^2}} \\
 &= \frac{\pi}{J}.
 \end{aligned}$$

3.C Quantifiers of mesh quality

For each triangle i of a mesh, we define the triangle skewness S_i as:

$$S_i = 1 - \frac{l_i^{\min}}{l_i^{\max}} \quad (3.23)$$

where l_i^{\min} and l_i^{\max} respectively depict the smallest and largest edge length of the triangle. Optimality ($S_i = 0$) is achieved only for equilateral triangles. The skewness is in fact a local quantifier of local triangle shape.

For each mesh node n we define the roughness R_n as:

$$R_n = 1 - \frac{\mathcal{A}^{\min}}{\mathcal{A}^{\max}} \quad (3.24)$$

where \mathcal{A}^{\max} and \mathcal{A}^{\min} respectively depict the surface areas of the biggest and smallest triangles containing node n .

3.D Measure of discrete correlation length

We consider a scalar quantity s defined at mesh nodes. We are interested in quantifying the spatial variation of s . For that we compute the discrete correlation function of s .

For each pair of node $(i, j) \in \mathcal{N}^2$ (\mathcal{N} is the set of all nodes), we define the topological distance $d(i, j)$ as the minimal number of triangle edges forming a path between nodes i and j (Fig. 3.12(a)). This distance allows to define the topological circle of center i and radius $L \in \mathbb{N}$ as:

$$\mathcal{C}(i, L) = \{j \in \mathcal{N} | d(i, j) = L\}. \quad (3.25)$$

(Fig. 3.12(a)).

We define the discrete autocorrelation function of s as:

$$a_s(L) = \frac{1}{\#\mathcal{N}} \sum_{i \in \mathcal{N}} \sum_{j \in \mathcal{C}(i, L)} \frac{(s_i - \bar{s})(s_j - \bar{s})}{\#\mathcal{C}(i, L)} \quad \text{with} \quad \bar{s} = \frac{1}{\#\mathcal{N}} \sum_{i \in \mathcal{N}} s_i. \quad (3.26)$$

This extends the classical definition of the autocorrelation function to a mesh. We define the discrete correlation length of s by performing an exponential fit on the discrete values $a_s(L)$, which provides a characteristic decay distance (Fig. 3.12(b)).

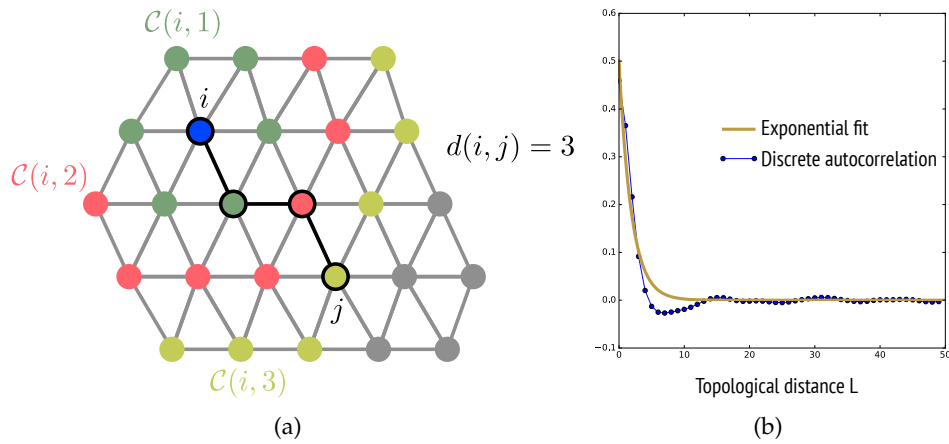


Figure 3.12: **(a)** Topological distance and circles. **(b)** Discrete autocorrelation function and exponential fit (example).

Chapter 4

Application: anisotropic growth of leaves

Foreword

This chapter corresponds to the work here referred as Zhao et al. (2019). An official preprint is available at www.biorxiv.org/content/10.1101/604710v1. I am third amongst four equally-contributing first authors of this work (F. Zhao, F. Du, myself and L. Zhou). I have conceived the 3D modeling strategy and the shape diagram introduced hereafter. I implemented and carried out the 3D simulations, and produced the related supplementary materials and figures, based on the 3D growth model detailed in Chapter 3. To generate meshes, I have implemented a semi-automated method for generating 3D alveolar structures (detailed in Ali et al., 2019, and in Section 4.A.5). Besides, I intensively took part of the discussions, and to the collaborative writing of the manuscript.

4.1 Abstract

Many plant species have thin leaf blades, which is an important adaptation that optimizes the exchanges with the environment. Here, we provide evidence that their three-dimensional geometry is governed by microtubule alignment along mechanical stress patterns in internal walls. Depending on the primary shape of the primordium, this process has the potential to amplify an initial degree of flatness, or promote the formation of nearly axisymmetric, mostly elongating organs, such as stems and roots.

This mechanism may explain leaf evolution from branches, which is alternative to Zimmermann's influential, but widely questioned, *telome* theory.

4.2 Main text

The formation of thin leaf lamina in plants is an important adaptation that optimizes vital processes, including photosynthesis, transpiration and respiration (Maugarny-Calès and Laufs, 2018). While the regulatory genetic network controlling leaf polarity has been well characterized (Kuhlemeier and Timmermans, 2016), comparatively little is known on how such a thin structure mechanically arises and maintains itself during development. We addressed this issue by combining computational modeling and a three-dimensional (3D) experimental analysis of leaf morphogenesis in two species (*Arabidopsis thaliana* and tomato, *Solanum lycopersicum*). Various leaf types (rosette and cauline leaves, cotyledons and sepals) were analyzed.

Primordia of leaves and leaf-like organs initiate from apical meristems, as rounded, slightly asymmetric bulges (Fig. 4.9c and d). Starting from a ratio of blade width (in the mediolateral axis) to thickness (in the dorsoventral axis) between 1.5 and 2, the leaf and sepal primordia mainly expand in two dimensions, forming a thin lamina with ratios of 10-12 in sepals and even higher in leaves (Poethig and Sussex, 1985) (Fig. 4.1a, d and Fig. 4.9). Growth directions largely rely on the orientation of the cellulose microfibrils in the cell walls (Ali et al., 2014, Sampathkumar et al., 2014a), which depends on the organization of the cortical microtubule (CMT) arrays guiding the cellulose synthase complexes (Paredes et al., 2006).

To investigate the role of CMTs in leaf development, we first characterized CMT arrangements using immunostaining and in vivo confocal imaging (Fig. 4.1b and e to j). CMT behavior along the inner and outer periclinal walls was highly dynamic. In very young growing sepals at stage 3/4, these CMTs transiently showed some degree of anisotropy (Fig. 4.1e, f), which decreased significantly (from 30% to 10%) early in development (Fig. 4.1h-j; see also Hervieux et al., 2016). Similarly, the very young leaf also transiently showed aligned CMTs along the outer periclinal walls, which became more disorganized afterwards (Fig. 4.10a to d).

A very different behavior was found along most of the anticlinal walls. Here CMTs were mainly oriented perpendicular to the surface in *Arabidopsis* cotyledons, leaves and sepals, as well as in tomato leaves (Fig. 4.1b-j, Fig. 4.10e and Fig. 4.11a, b). Staining of cellulose confirmed that this coincided with the main microfibril orientation in

these walls (Fig. 4.1c and Fig. 4.11c), while the cellulose synthase-associated proteins followed anticlinal paths along the CMTs (Fig. 4.11d-i).

To further evaluate the role of the CMTs in leaf development, we treated primordia with the CMT-depolymerizing drug *oryzalin*, at concentrations where they continued to grow. After treatment, the width to thickness ratio did not increase, in contrast to the untreated controls (Fig. 4.1k-p and Fig. 4.13). Outgrowing leaves and sepals were thicker, while lateral expansion was compromised. When cells continued to divide, division plane alignment became randomized. This shows that CMTs arrays are crucial for asymmetric leaf expansion.

How do these heterogeneous and dynamic CMT arrangements on anticlinal and periclinal membranes emerge? In plants, turgor pressure and differential growth both generate mechanical stresses within the cell walls (Peters and Tomos, 1996). It has been proposed that these stresses serve as a regulatory cue for cellular growth (Ali et al., 2014, Sampathkumar et al., 2014a). Indeed, CMTs often align along the axis of maximal tension (Hamant et al., 2008, Landrein and Hamant, 2013b). This in turn would lead to the CMT-guided deposition of cellulose microfibrils (Landrein and Hamant, 2013b) and wall reinforcement restricting growth in this orientation of maximal tension (Fig. 4.11d-i).

We first investigated if this so-called stress feedback mechanism could provide, on theoretical grounds, a plausible scenario for leaf morphogenesis. To this end, we developed a computational modeling approach. The models of leaf development proposed in the literature – (e.g. Hervieux et al., 2016, Kuchen et al., 2012, Qi et al., 2017) – are in 2 dimensions. Therefore, out-of-plane walls are not taken into account, although they could significantly impact the mechanics of the system. To alleviate this limitation, we developed a 3D finite element (FE), multicellular model (adapted from Boudon et al., 2015, Oliveri et al., 2019, see also Appendix 4.A), to analyze the effect of mechanical feedback at the level of the entire growing organ. The effect of cell division, not taken into account in these simulations, was considered to be negligible as the simulations were only carried out over short time periods.

Incipient leaves were represented as ellipsoidal alveolar structures (composed of 800 cells) under steady and uniform pressure (see Appendix 4.A). We used ellipsoids of initial aspect ratios comparable to that of young primordia (Fig. 4.2). To account for the differences in thickness between outer and inner walls observed *in vivo* (Fig. 4.14), the outer walls were made 3 times stiffer than the inner walls in the model.

Inside these structures the dominant wall forces were in the dorsoventral direction (*i.e.* along shortest axis of the ellipsoid). By contrast, at the outer surface, wall strain and

stress were dominantly mediolateral (*i.e.* along the second axis of the ellipsoid). Accordingly, the virtual structure evolved towards a spherical shape if no mechanical feedback was introduced (simulation 1: Fig. 4.2b, d), which echoes the results obtained *in vivo* using oryzalin treatment. Conversely, when the stress feedback was active throughout the entire 3D tissue, the structure grew longer (and slightly wider), showing that a stress feedback has the potential to promote anisotropic expansion (simulation 2: Fig. 4.2b, d, outer and inner feedback active). In line with the experimental evidence, these simulations showed CMT alignments along the dorsoventral axis on anticlinal walls (Fig. 4.2c, left panel), restricting growth in this direction.

However, simulations where the feedback was active throughout the entire tissue, systematically predicted a mediolateral alignment of CMT arrays along the stiffer outer walls (Fig. 4.2c, left panel), which is not consistently seen *in vivo*. In addition, we observed, *in silico*, an emergent loss of cohesion in CMT alignment between layers. Indeed, CMTs along the outer and inner periclinal walls were oriented perpendicularly (Fig. 4.15). This peculiar effect, also not seen *in vivo*, probably results from the apical-basal growth of the outer wall (itself prescribed by the mediolateral cellulose orientation), which in turn is actively resisted by the stress responsive inner tissue.

To rule out this effect, we next performed simulations where mechanical feedback was this time active on inner walls only. This scenario follows our *in vivo* observations showing disorganized CMTs on the outer periclinal walls of growing leaves and sepals (Fig. 4.1h-j, Fig. 4.10b-d). Here, simulations led to further amplification of flatness (simulation 4: Fig. 4.2b-d), while the predicted arrangements of CMTs along both outer and inner walls were qualitatively in line with the *in vivo* observations (Fig. 4.2c, right panel). This scenario, involving an uncoupling between CMTs in inner and outer walls, seems therefore more plausible than the previous one. Note that, by contrast, activating mechanical feedback on outer walls only, resulted in reduced asymmetry as the virtual organ developed towards an axisymmetric elongated shape (simulation 3: Fig. 4.2b and d).

A number of observations on *katanin* (*ktn*) mutants provided proof for such an uncoupling between outer and inner walls. KTN is involved in CMT alignment and its mutation leads to the formation of isotropic CMT arrays (Bouquin et al., 2003, Uyttewaal et al., 2012). Different from wild type leaves, the CMTs on the outer periclinal membranes were systematically more isotropic in the *ktn* mutants *bot* and *lue1* (Fig. 4.3a-c). However, the degree of anisotropy of anticlinal CMTs was not affected by the mutation (Fig. 4.3d, e). This indicates that in the mutants the CMTs on the outer wall never align with the predicted stress patterns, in contrast to the inner, anticlinal walls. Consistent with our simulations, the mutant leaf and sepal blade were relatively wider, while

maintaining a thickness at wild type levels (Fig. 4.3g, h). In summary, the results so far suggest a scenario, where CMTs systematically align along predicted stress patterns in internal walls. While this alignment guarantees leaf flatness, the degree of feedback on the outer cell wall is variable, and accounts for leaf blade width.

The previous simulations were initialized with a relatively flat ellipsoid. We also investigated the response of the system, starting from prolate spheroids only slightly perturbed in their degree of axisymmetry. This showed that the degree of flattening not only depended on the activation of the feedback itself, but also on the initial degree of shape asymmetry. Indeed, *in silico*, nearly spheroidal structures mainly grew in the apical-basal direction, resulting in the formation of elongated, finger-like shapes (simulation 5: Fig. 4.2b-d). An axisymmetric structure would maintain itself as such, as the feedback mechanism on inner walls, is on its own not sufficient to break axisymmetry (Fig. 4.2b).

We next tested experimentally the predicted link between initial primordium flatness and final flatness *in vivo*, using the sepal, which is easily accessible for observation. Leaf margin genes, such as *WOX1* and *PRS/WOX3*, are expressed in the lateral and middle domains (Fig. 4.16a, b), and are essential for setting up initial asymmetry (Guan et al., 2017, Nakata et al., 2012, Nardmann and Werr, 2013). Accordingly, plants with a double knock-out in both genes showed slightly narrower leaves and a clear reduction in sepal width (Fig. 4.3b and Fig. 4.16c, d). As the primordia of these mutants are still somewhat flattened, we would predict that reducing the initial mechanical feedback on the outer periclinal walls would partially rescue the narrow sepal phenotype. To this end, we introduced *bot* into a *wox1 prs* background, and found the width to thickness ratio of the sepals dramatically increased (Fig. 4.4a-d).

We tested a further reduction in primordium width/thickness ratio by combining *wox1 prs* with a third mutation, *asymmetric leaves2 (as2)*, which has elongated and (nearly) axisymmetric sepals and leaves Nakata et al. (2012) (Fig. 4.4e, f). Although still set up as a slightly flattened structure, the primordia soon became almost axisymmetric. They then mainly grew in the apical-basal direction (Fig. 4.3e and Fig. 4.17). The GFP-MBD marker revealed highly anisotropic CMT arrays on the anticlinal walls of sepals (not shown) and leaves (Fig. 4.4l). Importantly, along periclinal walls in *wox1 prs as2*, however, different arrangements were found, i.e. more isotropic in leaves and highly anisotropic in sepals (Fig. 4.4i and k). In other words, there is no specific CMT arrangement on the outer walls that correlates with the elongated shape of these organs. Therefore, axisymmetric or nearly axisymmetric shapes only seem to require CMT alignment

along anticlinal walls. Consistent with theory, *bot* was not able to restore axisymmetric shapes in this background: sepals were shorter and thicker, but remained close to axisymmetric in the *vox1 prs as2 bot* quadruple mutant (Fig. 4.4g).

The results in *Arabidopsis* were further confirmed using microsurgery in tomato. Isolation of an incipient leaf primordium from the meristem resulted in compromised WOX expression, leaf margin formation, and flattening (Shi et al., 2017). Both CMT arrays and cellulose microfibrils showed anticlinal arrangement in these axisymmetric leaves (Fig. 4.4m-p).

In conclusion, we have identified a conserved mechanism involving the coordinated behavior of the cytoskeleton in response to mechanical stress in internal, anticlinal walls. Although the precise mechanism behind this behavior remains elusive (Landrein and Hamant, 2013b), we suggest that a stress feedback has both the potential to amplify bilateral asymmetry during leaf development and to promote the elongation of stem-like organs, such as roots. This is a robust property, which is reproduced in our model with a minimum of hypotheses. In addition to the reinforcement of anticlinal walls along stress patterns, leaf flatness could in principle be further enhanced through cell division plane alignment in the same direction. It is known that cells often divide in a plane parallel to the microtubule interphase array (Ehrhardt and Shaw, 2006, Wasteneys, 2002). Accordingly, we observed that division planes were mostly perpendicular to the plane of the leaf blade (Fig. 4.1k and m, Fig. 4.12 and Fig. 4.13a and b). Such anticlinal walls should in principle further increase the resistance of the tissue to thickening, and thus both cellulose deposition and the orientation of new cross walls would contribute synergistically to the final leaf shape.

The amplification of asymmetry potentially provides a parsimonious explanation for leaf evolution. The widely accepted Zimmermann's telome theory proposes that a stem (telome) evolved into a thin leaf through series of shape transformations, which lack plausible molecular evidence (Beerling and Fleming, 2007). According to our model, once asymmetry is established in a primordium, the CMT-mediated mechanical feedback would amplify the asymmetry to form a plenary leaf blade. Initial symmetry breaking can result from the asymmetric gene expression patterns at the shoot meristem, and likely involves asymmetric patterns of cell wall stiffness and expansion during early stages of development, in particular at the leaf margins (Guan et al., 2017, Nakata et al., 2012, Qi et al., 2017).

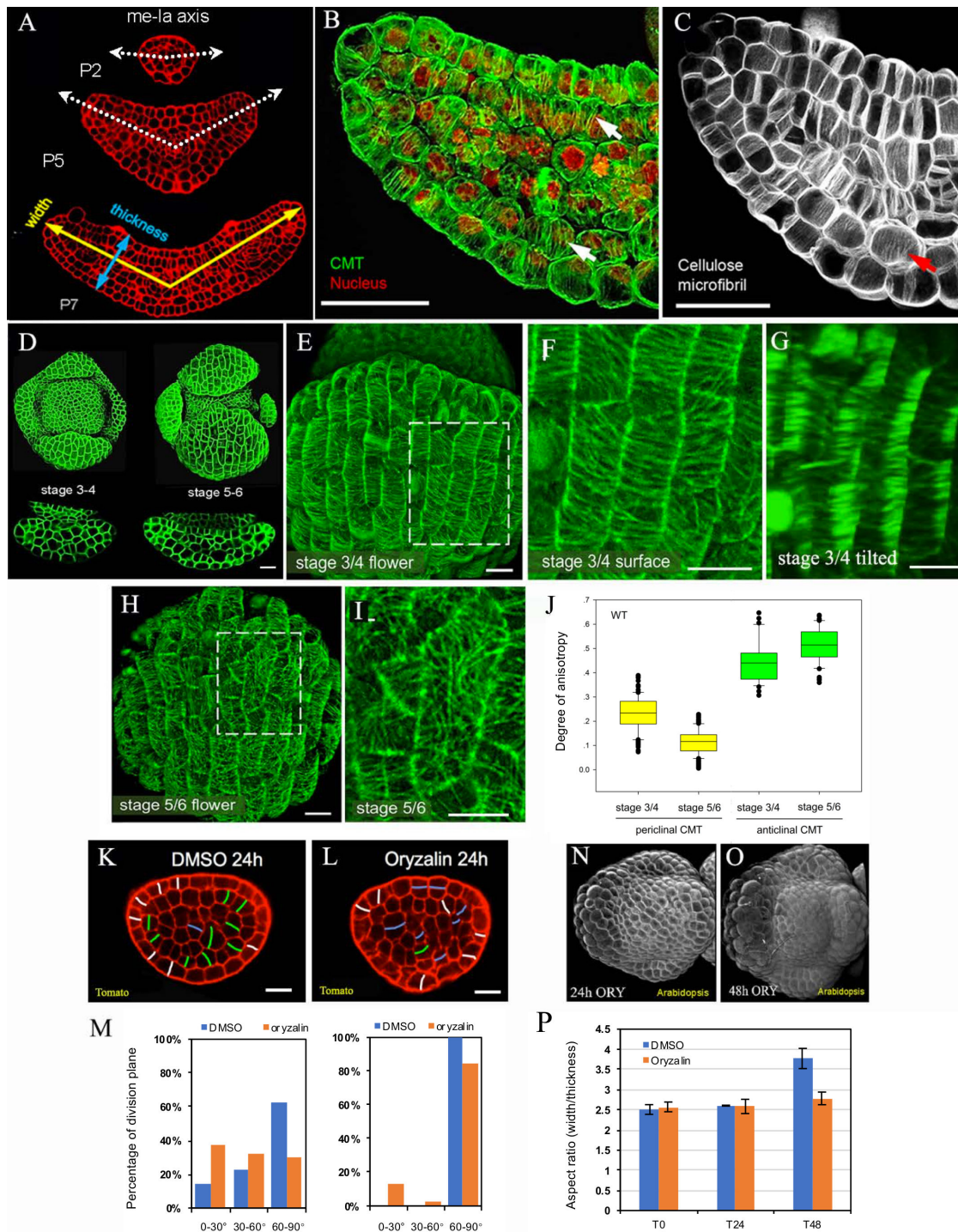


Figure 4.1: Continued on next page.

4.3 Acknowledgments

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Figure 4.1: Shape, CMT organization, and cell division in developing leaves and sepals. (a) Cross sections of *Arabidopsis* leaf primordia showing highly anisotropic growth. (b) CMT organization by immunostaining (green) with nuclei stained by DAPI (red) (c) CMFs stained by Direct Red 23 (white) in *Arabidopsis* rosette leaf primordia. (d) Overview of the same flower bud and cross section through the abaxial sepal at stages 3/4 and 5/6. (e-i) flowers sepals expressing GFP-MBD. (e) Overview of sepal at stage 3/4, inset indicates detail given in (f) showing anisotropic CMTs at the outer wall. (g) Same image stack as (f) but tilted to show highly anisotropic anticlinal CMTs (arrows). (h) Overview of sepal at stage 5/6, inset indicates detail given in (i) to show isotropic CMTs. (j) Quantification of CMTs on periclinal and anticlinal walls in sepals using Fibriltool (Boudaoud et al., 2014), showing differences in the degree of anisotropy along anticlinal (N=32 walls from 4 stage 3/4 sepals; N=52 walls from 5 stage 5/6 sepals) and periclinal (outer) (N=100 cells from 4 stage 3/4 sepals; N=207 cells from 5 stage 5/6 sepals) walls during sepal development. (k-m) Cell division pattern by mPS-PI staining in optical cross sections of tomato P3 treated with DMSO (k) or oryzalin (n) White, divisions perpendicular to the epidermis; blue, divisions parallel (angle $< 30^\circ$) to mediolateral axis in inner cells or to the epidermis; green, other divisions ($30^\circ \leq \text{angle} \leq 90^\circ$). Statistics are provided in (m). (n-p) Effect of oryzalin treatment on *Arabidopsis* sepal development after 24h (n) and 48h (o). (p) Quantification of width/thickness ratios. Treated sepals do not flatten (N=3 biological repeats). Scale bars, 20 μm in (a-c), 20 μm in (d), 10 μm (e-i).

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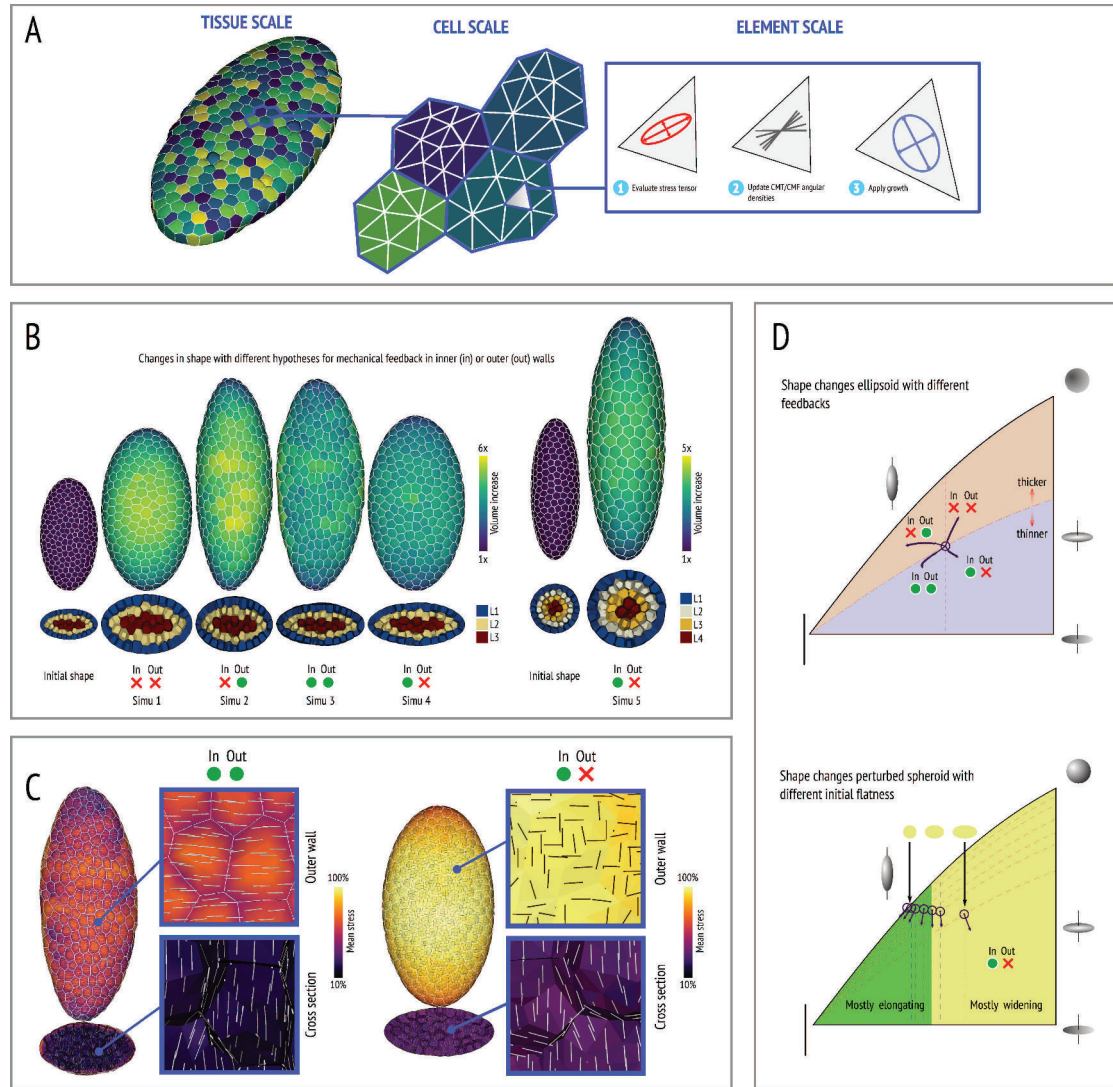


Figure 4.2: Computational modeling. (a) Overview of the 3D model. Virtual tissues with outer and inner cells are composed of triangular FEs. At each time step, the element stiffness is updated in function of the stress tensor (Chapter 3). (b) Outcome of five different scenarios (result after 20 time steps). Starting from a flattened ellipsoid (simulation 1-4) different levels of flatness amplification can be achieved depending on whether the feedback is active (green dot) or inactive (red cross) on outer and/or inner walls. Without feedback (simulation 1) the structure becomes thicker, with feedback everywhere (simulation 2) the structure becomes slightly flatter, with feedback on the outside only (simulation 3) the structure will become thicker and move towards a cylindrical shape. Maximal flattening is obtained with feedback on inner walls only (simulation 4). A prolate spheroid (simulation 5) will remain axisymmetric with the same feedback on inner walls only. (c) Predicted CMT orientations in simulation 2 (upper panel) and 4 (lower panel). Both simulations predict anisotropic CMTs/CMFs along anticlinal walls as observed *in vivo* (see white line segments on cross-sections). Simulation 2 leads to highly anisotropic CMTs/CMFs on outer walls (white line segments on outer wall), which is not always observed *in vivo*. (d) Top diagram: ellipsoid shape changes. These can be represented as respectively points and trajectories on a 2D diagram (Appendix 4.B). Feedback in the inner tissues causes flattening (trajectories below the dotted line). Bottom diagram: in (near) axisymmetric structures elongation dominates (trajectories in green area).

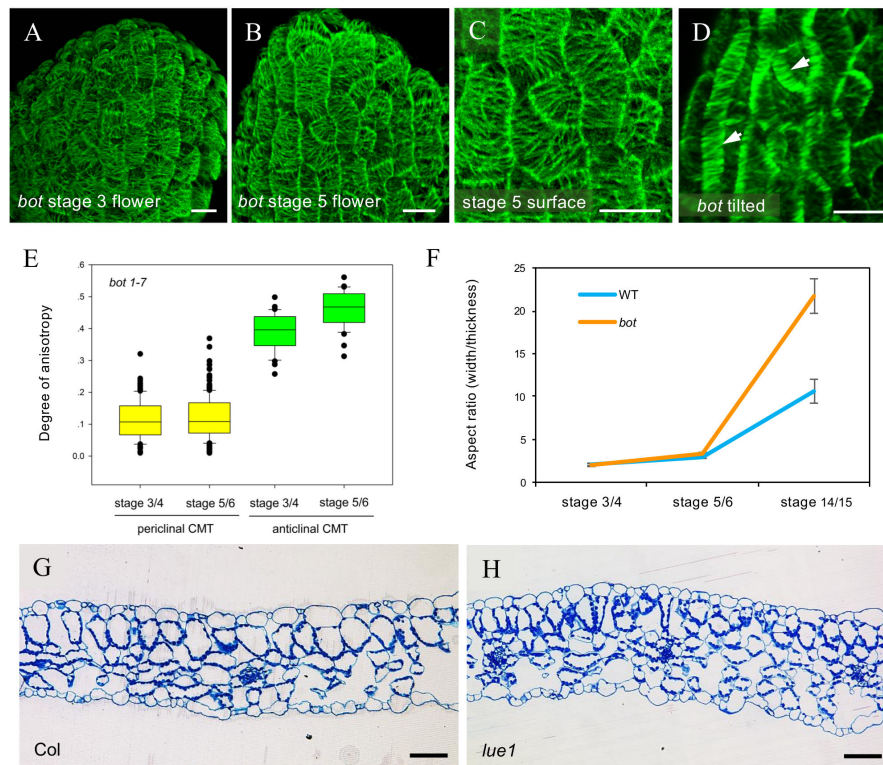
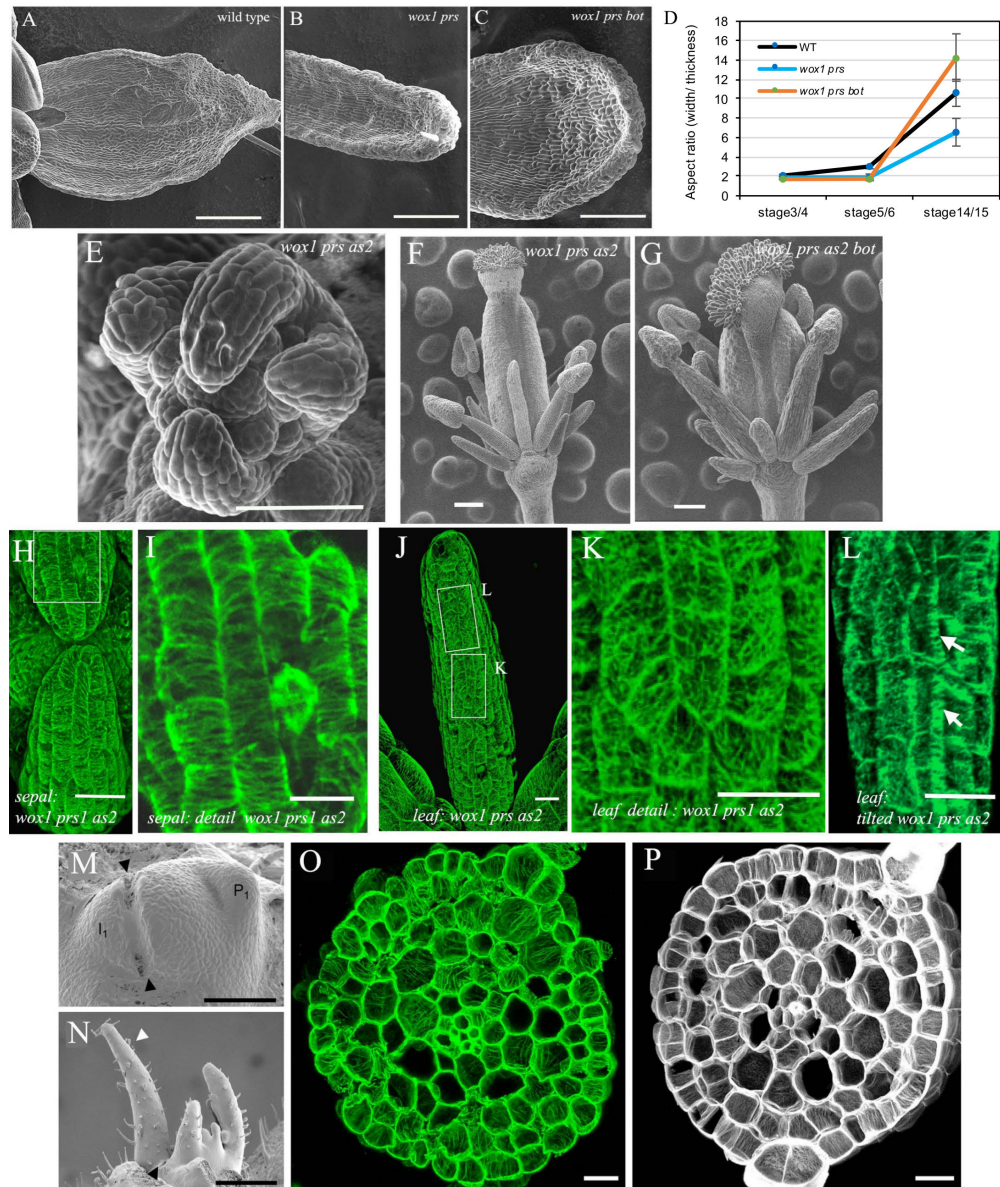


Figure 4.3: CMT arrangements in *ktn* mutant.(a) Overview of stage 3 *bot* flower bud expressing GFP-MBD showing more or less isotropic arrays. (b) Overview of stage 5 *bot* flower bud with isotropic CMTs. (c) Detail showing random orientation of periclinal CMTs in sepal in (b). (d) Tilted detail of (b). Anticlinal CMTs remain highly anisotropic in *bot* (indicated by arrows). (e) Quantification of microtubule orientations using Fibril-tool. Wild type sepals at stage 3/4 have a higher degree of anisotropy on their outer periclinal walls than at stage 5/6, while in *bot* mutants anisotropy is low from early stages onwards. By contrast CMTs on anticlinal walls remain highly anisotropic throughout development. N=100 cells from 4 stage 3/4 sepals and N=255 cells from 7 stage 5/6 sepals for periclinal analysis; N=31 walls from 3 stage 3/4 sepals and N=36 walls from 4 stage 5/6 sepals for anticlinal CMT analysis. (f) Quantification of width/thickness ratios in Col-0 and *bot* sepals. N=10 WT and 7 *bot* sepals at stage 3/4. N=9 WT and 11 *bot* sepals at stage 5/6. N=11 WT and 8 *bot* sepals at stage 14/15. (g,h) Cross sections of mature leaves of Col-0 wild type (g) and *lue1* (h). Scale bars, 10 μ m in (a-d); 50 μ m in (g, h).



Appendices

4.A 3D growth model

Author's note

This section is of the supplementary materials associated with the present preprint. It here mostly serves the self-containment of this chapter and contains a number of redundancies with respect to Chapters 2 and 3. However, several modeling specificities are here introduced, motivated by the use of more complex structures.

4.A.1 General ideas

The tissue is modeled as a multicellular alveolar structure, each cell being described as a set of connected walls, which together form a 2D continuum, constantly loaded with a steady and uniform pressure. To model both the elastic and plastic effects occurring in the virtual organ, we adapted the model detailed in [Boudon et al. \(2015\)](#), which represents the growing tissue as a *morphoelastic* system ([Goriely and Ben Amar, 2007](#), [Rodriguez et al., 1994](#)). This consists in postulating that the total deformation gradient \mathbf{F} (from the initial to the current configuration) is the product of an irreversible (plastic) component \mathbf{F}_g and a reversible (elastic) component \mathbf{F}_e (the subscripts 'g' and 'e' respectively stand for *growth* and *elastic*):

$$\mathbf{F} = \mathbf{F}_e \cdot \mathbf{F}_g \quad (4.1)$$

(see Fig. 4.5). Constitutively, in the morphoelastic approach, we assume that the strain-energy density Ψ is a function of \mathbf{F}_e only. By contrast, the *growth tensor* \mathbf{F}_g defines a stress-free configuration. In so far as each individual region of the domain may grow independently from its neighboring, the growth tensor \mathbf{F}_g in general does not define a compatible configuration, *i.e.*, it is not the gradient of a continuous displacement field

(in contrast to \mathbf{F}). Compatibility is ensured by means of the elastic tensor \mathbf{F}_e , at the price of introducing mechanical stresses.

We take advantage of the fact that growth (that relies on cell physiological processes) occurs much slower than elastic relaxation. This allows us to treat the plastic and elastic regimes separately. Each step of the algorithm can be decomposed in two sub-operations (detailed next), that consist of (i) computing the static equilibrium (Section 4.A.2.1), which provides a corresponding value of \mathbf{F}_e ; and (ii) computing growth by incrementally modifying \mathbf{F}_g (Section 4.A.3).

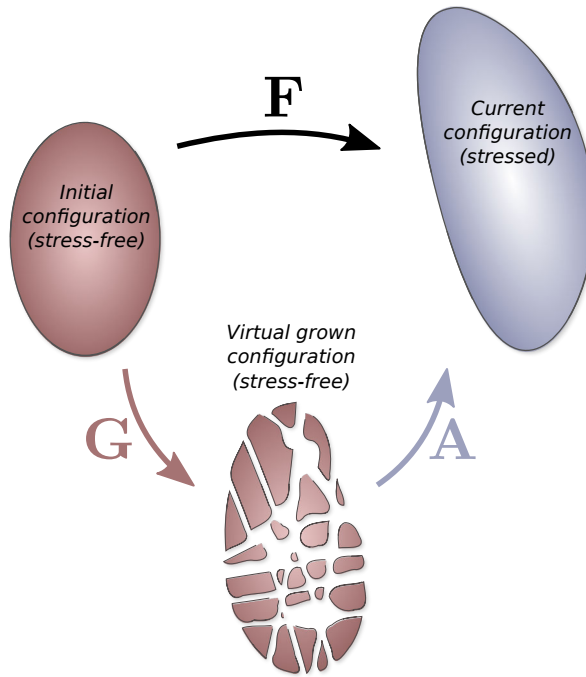


Figure 4.5: Morphoelastic description of growth kinematics. Cartoon of the multiplicative decomposition of the total deformation. The current configuration is obtained by composing the growth tensor \mathbf{F}_g that defines the stress-free state of the system (that is generally incompatible) and the elastic tensor \mathbf{F}_e that introduces mechanical stresses.

4.A.2 Elastic regime

4.A.2.1 Elastic model

We locally model the wall as a thin membrane under plane stress, described through the following strain-energy function:

$$\Psi(\mathbf{E}) = \frac{1}{2} \mathbf{E} : \mathbb{C}_w : \mathbf{E} \quad (4.2)$$

where operator \cdot is the tensor double-dot product; \mathbb{C}_w depicts the fourth order wall elasticity tensor; and

$$\mathbf{E} \stackrel{\text{def.}}{=} \frac{1}{2} (\mathbf{F}_e^T \cdot \mathbf{F}_e - \mathbf{I}) \quad (4.3)$$

is the *Green-Lagrangian* strain tensor associated with the elastic deformation (\mathbf{I} denotes the second order identity tensor).

To model the fibrous structure of the wall, the elasticity tensor \mathbb{C}_w is expressed as a function of the angular density of cellulose microfibrils, as detailed in [Oliveri et al. \(2019\)](#). We describe the CMF distribution on each element using the local (π -periodic) density function ρ (in practice, we only store a truncated Fourier decomposition of this distribution). The wall elasticity tensor is detailed using a Cox-like fibrous model [Cox \(1952\)](#), expressing \mathbb{C}_w as a function of the low-frequency Fourier coefficients of the CMF angular density. These coefficients are here noted $\{\rho_m\}_{m \in \mathbb{N}}$ and $\{\tilde{\rho}_m\}_{m \in \mathbb{N}}$, respectively representing the real and imaginary coefficients, and are given by:

$$\forall m \in \mathbb{N} : \quad (\rho_m, \tilde{\rho}_m) \stackrel{\text{def.}}{=} 2 (\Re(\hat{\rho}_m), -\Im(\hat{\rho}_m)) \quad \text{with} \quad \hat{\rho}_m \stackrel{\text{def.}}{=} \int_{\pi}^{\pi} \frac{d\theta}{\pi} \rho(\theta) e^{-2im\theta}. \quad (4.4)$$

where θ is and angle parameter in the orthonormal material basis $\{\mathbf{i}, \mathbf{j}\}$ associated with the triangle in its stress-free configuration (in which the stress, strain and elasticity tensors are expressed) The fiber-related stiffness is proportional to some fiber rigidity constant Y_f . As originally done in [Oliveri et al. \(2019\)](#), we also model the isotropic hydrogel embedding the fibers through an additional isotropic tensor, defined by its Poisson ratio ν and a reduced Young modulus Y (i.e. the actual Young's modulus divided by $1 - \nu^2$). The wall elasticity tensor is expressed in Voigt notation as:

$$[\mathbb{C}_w] = \underbrace{\frac{\pi Y_f \rho_0}{16} \begin{bmatrix} 3 + \frac{\rho_2 + 4\rho_1}{\rho_0} & 1 - \frac{\rho_2}{\rho_0} & \frac{2\tilde{\rho}_1 + \tilde{\rho}_2}{\rho_0} \\ 1 - \frac{\rho_2}{\rho_0} & 3 + \frac{\rho_2 - 4\rho_1}{\rho_0} & \frac{2\tilde{\rho}_1 - \tilde{\rho}_2}{\rho_0} \\ \frac{2\tilde{\rho}_1 + \tilde{\rho}_2}{\rho_0} & \frac{2\tilde{\rho}_1 - \tilde{\rho}_2}{\rho_0} & 1 - \frac{\rho_2}{\rho_0} \end{bmatrix}}_{\text{Microfibril elasticity matrix}} + \underbrace{Y \begin{bmatrix} 1 & \nu & 0 \\ \nu & 1 & 0 \\ 0 & 0 & \frac{1-\nu}{2} \end{bmatrix}}_{\text{Hydrogel elasticity matrix}}. \quad (4.5)$$

The values of Y and ν had little qualitative effect on the simulations (not shown). In the simulations presented here we used $\nu = 0.2$ – which is comparable to the values used in [Bozorg et al. \(2014\)](#), [Hamant et al. \(2008\)](#) – and $Y_f \rho_0 \sim 10^2 Y$, meaning that the mechanical contribution of the soft hydrogel is negligible in our study.

4.A.2.2 Numerical resolution of the static equilibrium

To compute the equilibrium in practice, we use triangular membrane finite elements equipped with \mathbb{P}_1 -Lagrangian shape functions ([Zienkiewicz et al., 2000](#)). Let \mathcal{N} and

\mathcal{T} be the respective sets of the mesh nodes and triangular elements. We iteratively integrate the equation of motion on nodes until the pressure forces \mathbf{f}_{ext} and reaction forces \mathbf{f}_{int} balance. This step is performed using the backward Euler method and the conjugate gradient to solve the intermediate linear systems.

At each sub-step, the nodal reaction force $\mathbf{f}_{\text{int}}^{(n)}$ (for each node $n \in \mathcal{N}$) is evaluated by differentiating the total potential strain energy U of the system with respect to node position $\mathbf{q}^{(n)}$:

$$\mathbf{f}_{\text{int}}^{(n)} = -\frac{\partial U}{\partial \mathbf{q}^{(n)}} = -\frac{\partial}{\partial \mathbf{q}^{(n)}} \left(\sum_{\tau \in \mathcal{T}} \epsilon^{(\tau)} \mathcal{S}^{(\tau)} \Psi^{(\tau)} \right) = -\sum_{\tau \in \mathcal{T}} \epsilon^{(\tau)} \mathcal{S}^{(\tau)} \frac{\partial \Psi^{(\tau)}}{\partial \mathbf{q}^{(n)}} \quad (4.6)$$

where $\mathcal{S}^{(\tau)}$ is the surface area of triangle $\tau \in \mathcal{T}$ (reference configuration); $\epsilon^{(\tau)}$ is the wall thickness at triangle τ (assumed to be constant in time, and either equal to ϵ_{in} for inner walls or $\epsilon_{\text{out}} = 3\epsilon_{\text{in}}$ for outer periclinal walls). The element-wise strain energy density $\Psi^{(\tau)}$ is obtained from Eq. (4.2):

$$\Psi^{(\tau)} \stackrel{\text{def.}}{=} \Psi \left(\mathbf{E}^{(\tau)} \right) = \frac{1}{2} \mathbf{E}^{(\tau)} : \mathbb{C}_{\text{w}}^{(\tau)} : \mathbf{E}^{(\tau)}. \quad (4.7)$$

In practice, Eq. (4.6) is evaluated by splitting $\partial \Psi^{(\tau)} / \partial \mathbf{q}^{(n)}$ in virtue of the chain rule:

$$\frac{\partial \Psi^{(\tau)}}{\partial \mathbf{q}^{(n)}} = \frac{\partial \Psi^{(\tau)}}{\partial \mathbf{E}^{(\tau)}} : \frac{\partial \mathbf{E}^{(\tau)}}{\partial \mathbf{F}_{\text{e}}^{(\tau)}} : \frac{\partial \mathbf{F}_{\text{e}}^{(\tau)}}{\partial \mathbf{q}^{(n)}}, \quad (4.8)$$

then using Eqs. 4.3 and 4.2 and the following expression:

$$\mathbf{F}_{\text{e}}^{(\tau)} = \sum_{n \in \mathcal{N}} \mathbf{q}^{(n)} \otimes \frac{\partial N^{(n)}}{\partial \mathbf{X}} \Big|_{\tau} \quad (4.9)$$

where \mathbf{X} depicts the 2D material coordinates in the grown configuration, and $N^{(n)}$ denotes the shape function associated with node n .

The external nodal loads (pressure forces) are given by:

$$\mathbf{f}_{\text{ext}}^{(n)} = \frac{1}{3} \sum_{\tau \in \mathcal{T}_n} s^{(\tau)} P^{(\tau)} \mathbf{n}^{(\tau)} \quad (4.10)$$

where \mathcal{T}_n is the set of the finite elements that contain node n ; $s^{(\tau)} = \mathcal{S}^{(\tau)} \det \mathbf{F}_{\text{e}}^{(\tau)}$ is the surface area of triangle τ in the current configuration; $\mathbf{n}^{(\tau)}$ is the normal to triangle τ ($\|\mathbf{n}^{(\tau)}\| = 1$); $P^{(\tau)}$ is the difference in pressure between both sides of element τ (the sign depends on the orientation of $\mathbf{n}^{(\tau)}$). We assume steady and uniform pressure P_{in} within the tissue. Hence, $P^{(\tau)}$ vanishes if triangle τ does not belong to an outer periclinal wall, and is equal to $P = P_{\text{in}} - P_0$ otherwise (P_0 being the outer atmospheric pressure).

4.A.3 Growth

4.A.3.1 Growth kinetics

To model growth we use the strain-based model developed in [Boudon et al. \(2015\)](#), expressing the *rate of growth* tensor \mathbf{L}_g as a piece-wise linear function of \mathbf{E} at equilibrium:

$$\mathbf{L}_g \stackrel{\text{def.}}{=} \frac{\partial \mathbf{F}_g}{\partial t} \cdot \mathbf{F}_g^{-1} = \Phi (\mathbf{E} - E_y \mathbf{I})_+ \quad (4.11)$$

The previous expression provides a multidimensional extension of Lockhart's one-dimensional model ([Lockhart, 1965](#)). Parameters Φ and E_y respectively depict the wall extensibility and yield threshold; $(\cdot)_+$ depicts the tensor ramp function. The latter is defined for any second order symmetric tensor \mathbf{T} as:

$$(\mathbf{T})_+ = \sum_k \max(0, T_k) \mathbf{T}_k \otimes \mathbf{T}_k \quad (4.12)$$

where T_k and \mathbf{T}_k are respectively the eigenvalues and corresponding normed eigenvectors of \mathbf{T} .

4.A.3.2 Numerical resolution of growth

Starting from a system at static equilibrium at t , Eq. (4.11) is integrated, for each triangle τ , using the forward Euler method with constant time step Δt :

$$\mathbf{F}_g^{(\tau)}(t + \Delta t) = \left(\mathbf{I} + \Delta t \mathbf{L}_g^{(\tau)}(t) \right) \cdot \mathbf{F}_g^{(\tau)}(t). \quad (4.13)$$

Since immediately after this operation, the node positions have not moved yet, Eq. (4.1) can be rewritten as:

$$\mathbf{F}^{(\tau)}(t) = \tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t) \cdot \mathbf{F}_g^{(\tau)}(t + \Delta t), \quad (4.14)$$

where $\tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t)$ defines the new elastic deformation, that compensates for the new growth tensor. In virtue of Eqs. 4.1, 4.13 and 4.14, $\tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t)$ can be computed from the previous elastic deformation $\mathbf{F}_e^{(\tau)}(t)$, according to:

$$\tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t) = \mathbf{F}_e^{(\tau)}(t) \cdot \left(\mathbf{I} + \Delta t \mathbf{L}_g^{(\tau)}(t) \right)^{-1} \quad (4.15)$$

($\mathbf{I} + \Delta t \mathbf{L}_g^{(\tau)}(t)$ is nonsingular). Eq. (4.15) expresses the idea that augmenting the plastic component amounts to relaxing some part of the elastic deformation by the same

amount. This allows to simulate growth without explicitly storing \mathbf{F}_g , in so far as only the dilation factor $\mathbf{I} + \Delta t \mathbf{L}_g^{(\tau)}(t)$ is needed to evaluate Eq. (4.15).

Note the presence of both times t and $t + \Delta t$ in Eq. (4.15). This reflects the fact that, *a priori*, $\tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t)$ does not satisfy mechanical equilibrium, hence the tilde notation. The deformation gradients $\tilde{\mathbf{F}}_e^{(\tau)}(t + \Delta t)$ and the current positions give the initial conditions to compute the next mechanical equilibrium (Section 4.A.2.2).

4.A.3.3 Growth-induced advection of the microfibrils

In principle, growth induces a change in the fiber density and orientation, which can be assimilated to an advective process. In fact, by virtue of mass conservation, the volumetric density of CMFs may decrease as the wall expands (if we disregard cellulose intake). Moreover, anisotropic growth modifies the orientation of each fiber, which rotates towards the direction of main deformation.

To model these effects, we assume that the deformation of the fibers is *affine*, namely that these deform according to the macroscopic medium. We update the microfibril angular density incrementally, for each triangle τ and at each growth step t (*N.B.*: for the sake of clarity, we drop the t and τ notations from now on). This is done by replacing the old fiber distribution ρ with the new one (noted $\tilde{\rho}$), which verifies:

$$\tilde{\rho}(\theta) \propto (1 - \alpha \cos(2(\theta - \vartheta)))^{-1} \rho(\Theta) \quad \text{and} \quad \int_{\pi} \tilde{\rho} = J^{-1} \int_{\pi} \rho \quad (4.16)$$

where $J \stackrel{\text{def.}}{=} \det(\mathbf{I} + \Delta t \mathbf{L}_g)$ depicts the incremental variation of volume; θ and Θ respectively parameterize fibers in the deformed (after incremental growth is applied) and initial (before incremental growth is applied) configurations; α and ϑ respectively measure the anisotropy and angle of the incremental growth.

In practice, this step is computed in the angle-domain, by first using the *Inverse Fast Fourier Transform* computed on the Fourier coefficients of ρ , which allows to compute Eq. (4.16); and then the *Fast Fourier Transform*, which provides the new Fourier coefficients of the microfibril distribution.

4.A.4 Stress feedback

To model the stress feedback, we express the dependency of cortical microtubules organization upon stress, and the CMT-guided cellulose deposition as detailed in [Olivieri et al. \(2019\)](#). Based on a kinetic model of CMT polymerization/depolymerization

(modulated by stress), the steady angular probability of presence ϕ of CMTs (at a given position) can be expressed as a function of stress and angle θ :

$$\phi(\theta) \propto \exp(\gamma \mathbf{S} : \boldsymbol{\theta} \otimes \boldsymbol{\theta}) \quad (4.17)$$

where γ is a sensitivity parameter; $\mathbf{S} \stackrel{\text{def}}{=} \partial(\epsilon\Psi)/\partial\mathbf{E}$ measures the *second Piola-Kirchhoff* (PK2) stress integrated over the wall thickness ϵ (i.e. the stress in the stress-free configuration of each triangle element, expressed in the 2D material coordinates); and $\boldsymbol{\theta} \stackrel{\text{def}}{=} \cos\theta \mathbf{i} + \sin\theta \mathbf{j}$.

Eq. (4.17) captures a CMT accumulation in the first direction of stress (that maximizes the argument of the exponential). Moreover, the sharpness of this alignment is a growing function of both stress anisotropy and stress amplitude (Oliveri et al., 2019).

We model the polymerization/depolymerization of cellulose fibers through (i) a deposition of cellulose in direction θ proportional to $\phi(\theta)$ (characterized by a kinetic constant k_ρ), coupled with (ii) a linear decay of cellulose (characterized by a kinetic constant k'_ρ):

$$\dot{\rho}(\theta) = k_\rho \phi(\theta) - k'_\rho \rho(\theta). \quad (4.18)$$

Eq. (4.18) is solved by the classical Runge-Kutta procedure (RK4) with constant time step size (the time step size Δt for growth and cellulose deposition are the same), assuming constant stress during the time step. At the beginning of the simulation, CMT and CMF distributions are isotropic and uniform, taking the trivial null stress steady solution of Eq. (4.18) as an initial condition for ρ :

$$\rho(\theta)|_{t=t_0} = \frac{k_\rho}{\pi k'_\rho}, \quad (4.19)$$

see Oliveri et al. (2019). After estimation of the first mechanical equilibrium, the CMF distribution is equated with the constant-stress steady solution of Eq. (4.18), that is proportional to ϕ . This allows for a coarse initialization of the CMF system at the very beginning of the simulation, which provides a more realistic initial condition than isotropic stiffness.

For better spatial regularity of the CMT density, we smooth the stress computed at each element over each cell interface, and use the smoothed expression of the stress in Eq. (4.17). Since the PK2 stress is given in the material coordinate system of each triangle, i.e. in the incompatible stress-free configuration, it cannot be directly used to perform a smoothing. In practice, this is performed by computing the stress in the current configuration of the material (Cauchy stress) from the PK2 stress \mathbf{S} . Then, we compute a mean stress over each cell interface (weighted by triangle area). The outcome of this

operation is generally of rank 3, as the interfaces may be slightly non planar. Therefore, the result is then re-projected onto the mesh triangles and pulled back to the reference configuration, providing a smoothed PK2 stress field.

4.A.5 Meshes

Each cell wall is composed of ~ 10 -20 triangular finite elements, and is common to two cells (unless it is an outer periclinal wall). Meshes are generated following the procedure detailed in [Ali et al. \(2019\)](#), itself based on the open source algorithm *DRACO-STEM* ([Cerutti et al., 2017](#)), which allows to build meshes dedicated to finite element modeling, from incoming cell segmentations, which are 3D images. Fig. 4.6(a) summarizes the pipeline. In order to generate abstract cellularized shapes such as ellipsoids (see Fig. 4.6(b)), a cell segmentation is artificially generated, in the form of a pseudo-random *centroidal Voronoi tessellation* (CVT) defined inside an input surface mesh (see Fig. 4.6(a)). A CVT is by definition a Voronoi tessellation wherein the centroid and seed of each region coincide. It is here generated using Lloyd’s algorithm ([Du et al., 1999](#)). The main benefit of a CVT, with respect to a more general Voronoi diagram, is that it allows more uniform cell volumes and shapes. This is a convenient property for us, as it allows to rule out the possible effects of cell shape and size variability on global morphogenesis. The CVT also builds cell layers that have a visually uniform thickness, which qualitatively resemble the actual cell layers seen in leaves.

4.A.6 Main parameters

In the simulations shown in the main text, we used the following parameters. For turgor pressure, we chose $P = 0.25$ MPa, which is of the order of magnitude of the values reported in [Beauzamy et al. \(2015\)](#). Meshes were scaled so that the average cell volume was equal to $V_{\text{cell}} \approx 5 \times 5 \times 5 \mu\text{m}^3$, which is close to what was measured in our experiments (each mesh contains 800 cells, total volume is $800V_{\text{cell}} \sim 10^5 \mu\text{m}^3$). The mean wall thickness was parameterized as $(\epsilon_{\text{out}} + \epsilon_{\text{in}}) / 2 = 0.25 \mu\text{m}$ (with $\epsilon_{\text{out}} = 3\epsilon_{\text{in}}$), which is close to the thickness measured experimentally in this work. The fiber rigidity is characterized by the Young modulus $Y_f k_\rho / k'_\rho \sim 10^2$ MPa ([Chanliaud et al., 2002](#)). The growth characteristic time was set as $\Phi^{-1} \approx 20$ min. For a strain of 5%, this roughly corresponds to an expansion speed of about 15% per hour (cell length doubles every 5h), which is close to the parameter used in [Lockhart \(1965\)](#).

Other parameters, which are less easily measurable, were set in a qualitative manner, in order to obtain a coherent simulation behavior: growth threshold $E_y = 1\%$, feedback

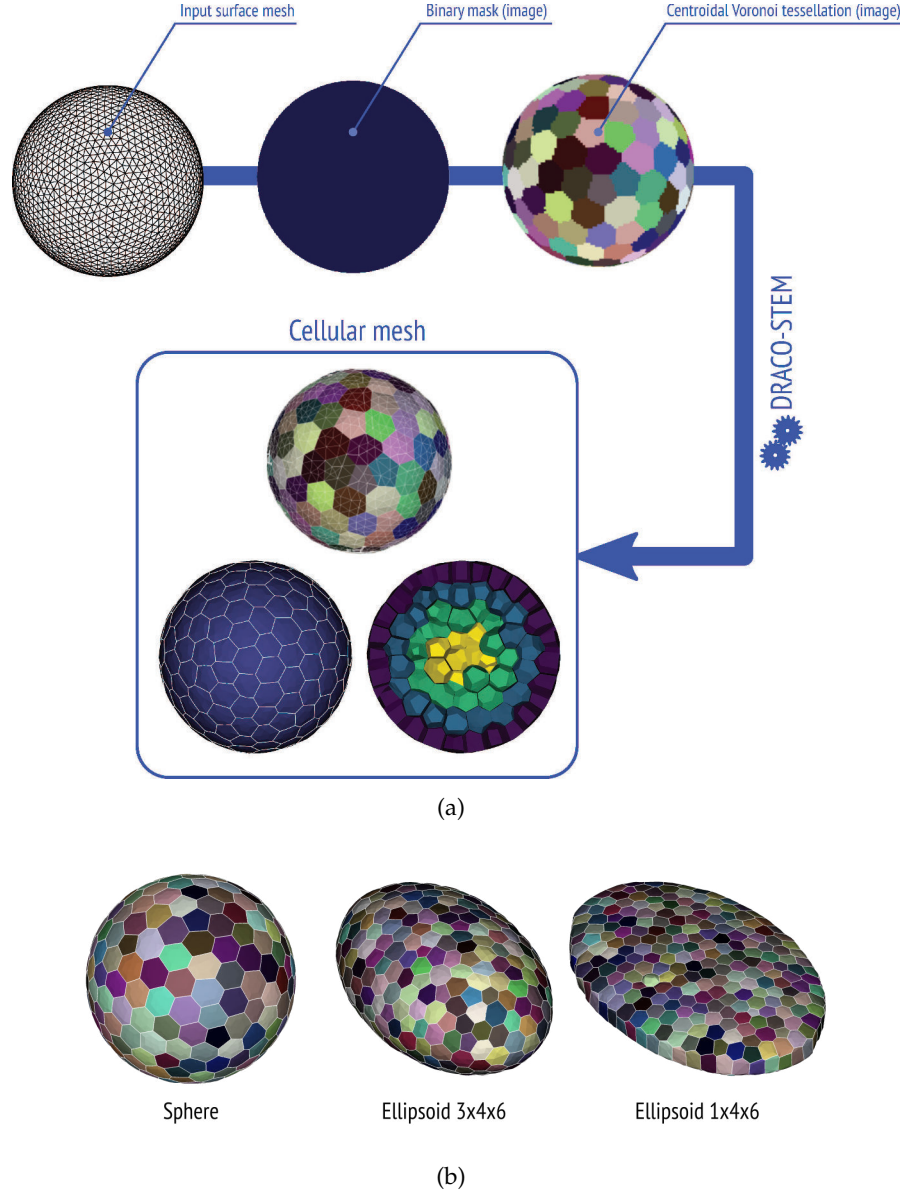


Figure 4.6: **(a)** Semi-automated generation of artificial multicellular tissue meshes: general pipeline. The algorithm takes a surface mesh as an input (here a spherical mesh). Note that the mesh quality at this stage is not important. A CVT is computed (Lloyd’s algorithm) within a binary mask, labeling the interior of the input mesh. The *DRACO-STEM* algorithm (Cerutti et al., 2017) allows to compute a final cellularized triangle tessellation from the incoming CVT. **(b)** Three examples of ellipsoidal cellular meshes.

sensitivity $\gamma = 30 \text{ MPa}^{-1} \cdot \mu\text{m}^{-1}$ (Section 4.A.8 provides a sensitivity analysis), cellulose deposition constant $k_p = 1 \text{ mol} \cdot \mu\text{m}^{-2} \cdot \text{h}^{-1}$, cellulose decay constant $k'_p = 0.5 \text{ h}^{-1}$.

4.A.7 Softwares

In-house *Python* code was implemented in order to simulate growth and stress feedback. To compute the static equilibrium, we employed the finite element method as implemented in the open source software *Sofa* (Faure et al., 2012) available at www.sofa-framework.org. Data structures (cellularized meshes) are implemented in the *Python* library *CellComplex* (gitlab.inria.fr/mosaic/cellcomplex). Visualization of 3D simulations was based on the open source software *TissueLab* (github.com/VirtualPlants/tissuelab). The meshing algorithm (Section 4.A.5) was initialized using meshes generated with the open source software *Blender* (www.blender.org). The *DRACO-STEM* tool (Cerutti et al., 2017) is available at gitlab.inria.fr/mosaic/draco_stem.

4.A.8 Sensitivity to initial position and coefficient γ

Fig. 4.7 provides a sensitivity analysis of the model with respect to initial shape (*i.e.*, initial position in the shape diagram) and feedback sensitivity γ , in the case where outer stress feedback is inhibited. This was performed by generating 7 different ellipsoidal meshes with equal volume and cell number, following the procedure detailed in Section 4.A.5. In this way, only the various aspect ratios varied between all the meshes. For null γ we robustly obtained trajectories heading towards a sphere (represented by blue trajectories in Fig. 4.7), which corresponds to the intuitive scenario where all walls are isotropic in stiffness. Higher γ generally resulted in flattening shapes (red and green trajectories in Fig. 4.7), for all the tested initial positions.

4.B Shape diagram

4.B.1 Construction

In order to monitor the change of a given growing shape, we track its principal dimensions over time, namely its length (λ_1), width (λ_2) and thickness (λ_3). In this work, we mostly focus on the various aspect ratios of the shape, and we disregard the volume of the structure, which, by contrast, characterizes the *size* of the organ. Hence, we can represent the shape at each time point through a normalized vector Λ defined as:

$$\Lambda_i = \frac{\lambda_i}{\sqrt{\sum_j \lambda_j^2}}. \quad (4.20)$$

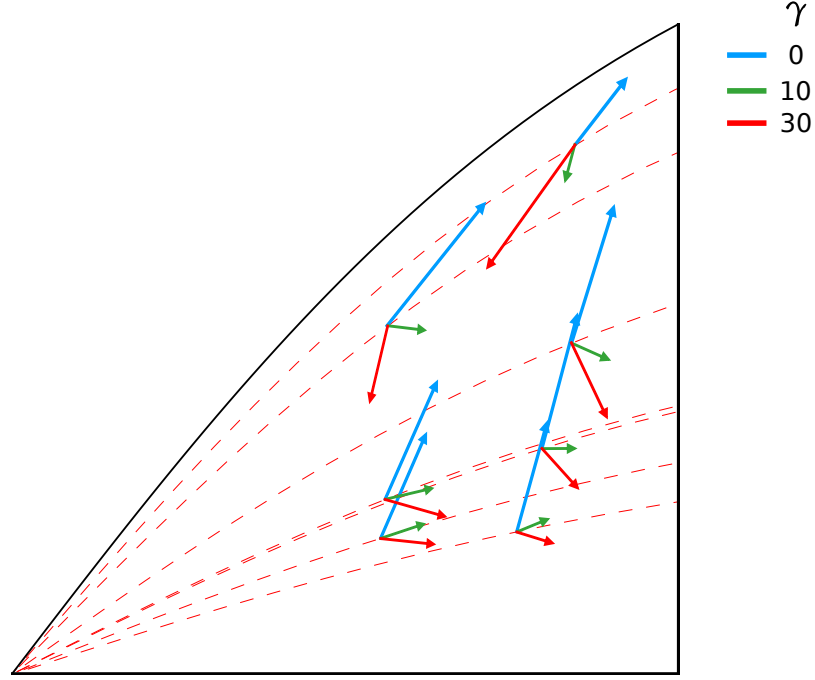


Figure 4.7: Sensitivity analysis with respect to initial shape and γ (in $\text{MPa}^{-1} \cdot \mu\text{m}^{-1}$), see Section 4.B. We used here $k'_\rho = 1 \text{ h}^{-1}$ and $\Phi^{-1} \approx 10 \text{ min}$.

In the sequel, we will note $\mathbf{\Lambda} = (L, w, T)$ the Λ -position of a given 3D shape. As $\mathbf{\Lambda}$ is of norm 1, it is equivalent to a point of the unit sphere (Fig. 4.8(a)). Hence, it can be equivalently represented using a longitude-latitude spherical parameterization (θ, φ) . Provided the natural constraint $L \geq w \geq T \geq 0$, the set of possible values for $\mathbf{\Lambda}$ is bounded by a spherical triangle defined by vertices $\mathbf{a} = (1, 0, 0)$, $\mathbf{b} = 1/\sqrt{2}(1, 1, 0)$ and $\mathbf{c} = 1/\sqrt{3}(1, 1, 1)$, or equivalently $(\theta_a, \varphi_a) = (0, 0)$, $(\theta_b, \varphi_b) = (\pi/4, 0)$ and $(\theta_c, \varphi_c) = (\pi/4, \pi/6)$ (Fig. 4.8(b)).

4.B.2 Interpretation

4.B.2.1 Vertices and edges

Vertex \mathbf{a} in Fig. 4.8(b) represents the infinitely prolate shapes (line). Vertex \mathbf{b} represents the shapes with null thickness, the other dimensions being equal (e.g. disc). The last vertex \mathbf{c} represents all the shapes having equal length, width and thickness (e.g. spheres). Note that vertices \mathbf{a} and \mathbf{b} represent degenerated shapes with null volume. Intermediately, edges \mathbf{ac} and \mathbf{bc} capture all possible shapes having two equal dimensions (prolate for \mathbf{ac} or oblate for \mathbf{bc}). Last edge \mathbf{ab} refers to all the degenerated shapes with null thickness.

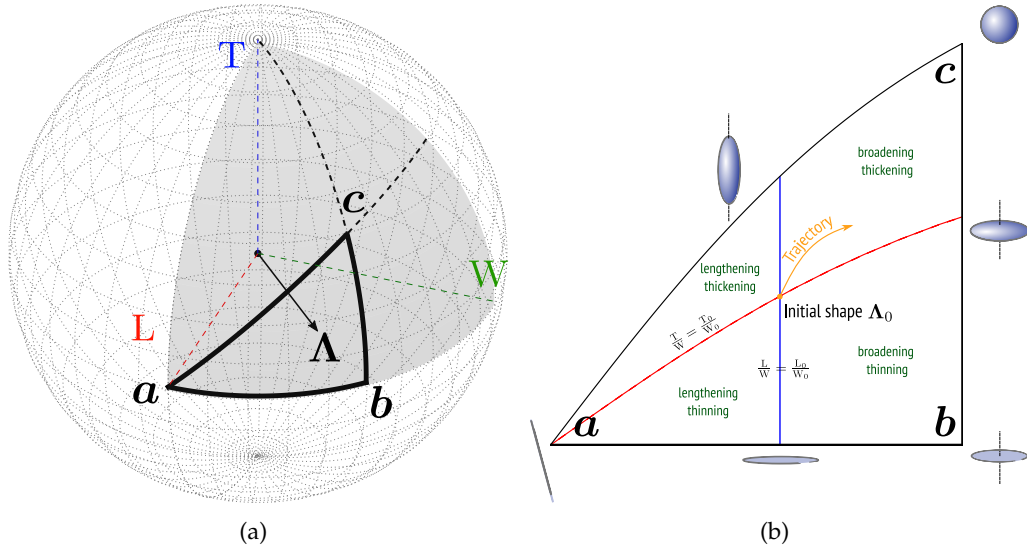


Figure 4.8: Morphospace. (a) 3D visualization of a shape vector Λ . (b) 2D spherical projection of the triangle morphospace.

4.B.2.2 Zones

Consider an initial shape represented by an initial Λ -position $\Lambda_0 = (L_0, W_0, T_0)$ within the triangle domain. As time progresses, the shape changes in time, and its 2D position moves in the domain, following a 2D trajectory. In order to interpret this trajectory in term of 3D growth, we may identify several useful lines which correspond to stereotypical growth behaviors (these lines are geodesics of the unit sphere). A trajectory that would follow the line defined by $L/W = L_0/W_0$ would represent a growth regime wherein only thickness (T) would vary relative to the other dimensions (blue vertical line on Fig. 4.8(b)). Similarly, line $T/W = T_0/W_0$ (red oblique line on Fig. 4.8(b)) captures regimes wherein only length would vary relative to the other dimensions (no change in aspect ratio would occur in the transverse cut). These two secant lines allow to divide the shape diagram in four sub-domains, which are associated with the initial position and correspond to four stereotypical expansion regimes (indicated by the green labels on Fig. 4.8(b)).

4.C Supplementary figures

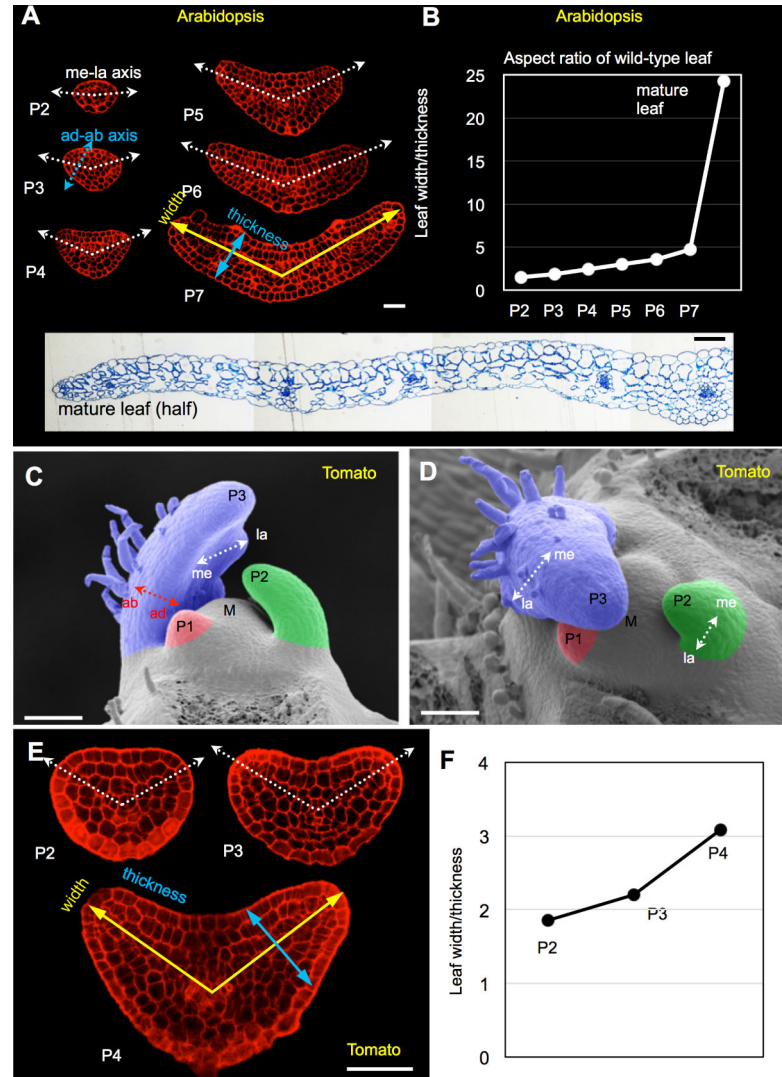


Figure 4.9: Anisotropic growth of Arabidopsis and tomato leaves. (a) Cross sections of Arabidopsis leaf primordia (P2-P7) showing that the growth of leaf primordia is highly anisotropic along the mediolateral axis (me-la axis, white) rather than the adaxial-abaxial axis (ad-ab axis, blue), which generates a planar form of mature leaves. (b) Quantification of width/thickness ratios in Arabidopsis leaves. (c, d) Scanning electron micrographs of a tomato shoot apex (c, side view; d, top view) show that the leaf primordia initiate surrounding the shoot apical meristem with an anisotropic growth more along the mediolateral axis (me-la) than along the adaxial-abaxial axis (ad-ab), resulting in a deformation from near symmetric (P1) to asymmetric (P2/P3) shape. P1 (red), youngest leaf primordium; P2 (green), second youngest leaf primordium; P3 (blue), third youngest leaf primordium; M, shoot apical meristem. (e) Cross sections of tomato leaf primordia (P2-P4) showing that the growth of leaf primordia is anisotropic along the mediolateral axis (white dash line). (f) Quantification of width/thickness ratios in tomato leaves. Leaf width is defined as twice the mediolateral axis connecting the centre to the farthestmost points on the outline of a leaf cross-section. Scale bar, 20 μm (white) and 50 μm (black) in (a); 100 μm in (c, d); 50 μm in (e).

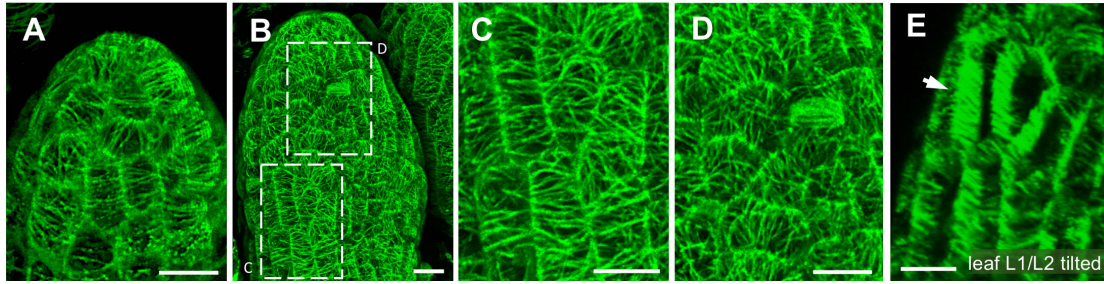


Figure 4.10: **CMT organization in developing Arabidopsis leaves.** (a) Young leaf primordium expressing GFP-MBD, top view, showing anisotropic CMTs . (b-d) Slightly older leaf showing different CMT arrangements. Boxed areas detailed in (c) (showing cells with anisotropic CMT arrays) and (d) (cells with isotropic CMT arrays). (e) Tilted version of (c), showing anticlinally oriented CMTs (arrow). Scale bars, 10 μ m.

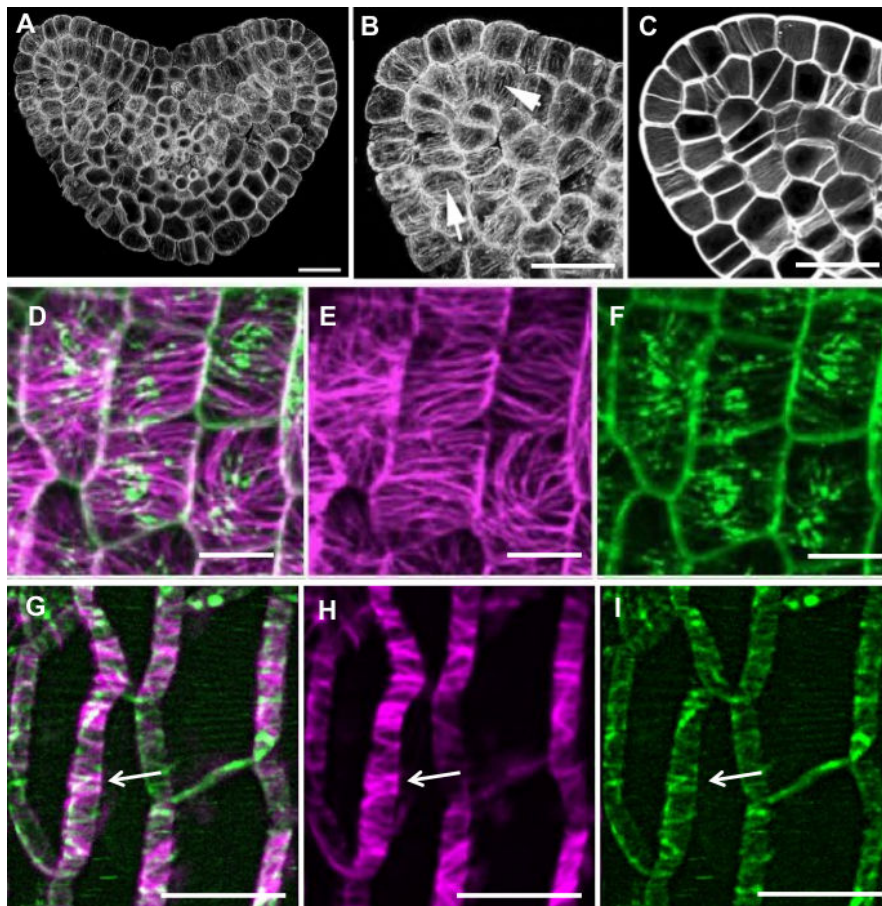


Figure 4.11: **Organization of anticlinal CMTs and CMFs in tomato and Arabidopsis.** (a-c) The organization of CMTs in the cross section of a tomato P3 by immunostaining with tubulin antibody. (a) overview, (b) shows the magnification of a part of (a), Arrow heads show CMTs. (c) The orientation of CMFs in the cross section of a tomato P3 stained by Direct Red 23 dye. (d-i) Live imaging of 3xYFP-labelled Cellulose Synthase Interacting 1 (CSI1/ POM2, green) in mCherry-TUA5 (pink) background sepals. (d-f) average projections to reveal CSI trajectories along microtubules on periclinal surface membrane. (g-i) show trajectories along anticlinal walls. Note anticlinal trajectories of CSI (arrows). Imaging by z-stacking (0.3 μ m steps) and time-lapsing (1 min intervals) for 20 min. Scale bars, 20 μ m in (a-c), 10 μ m in (d-i).

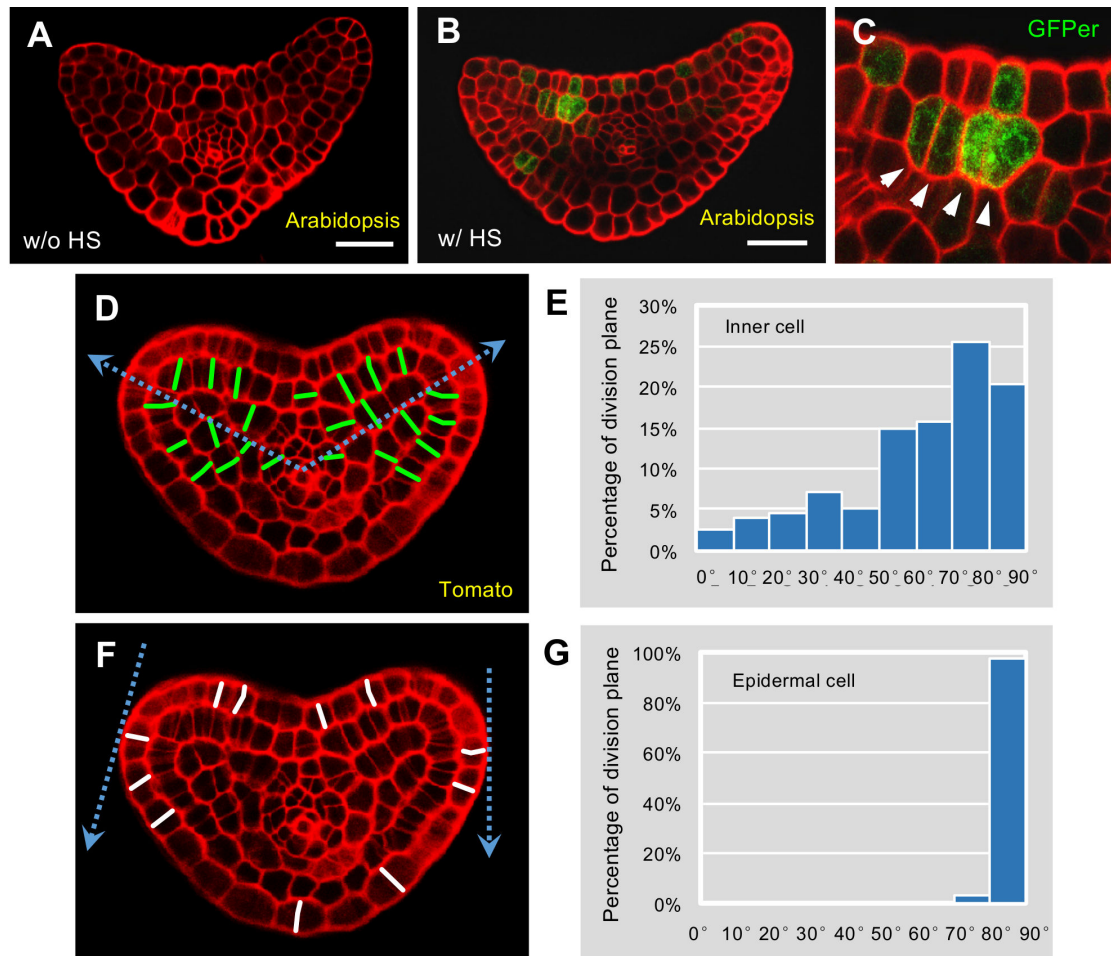


Figure 4.12: Division orientations in inner and epidermal cells of leaf primordia. (a-c) Cell lineage tracing analysis in Arabidopsis leaf primordia using a heat-shock induced Cre-loxP system. No endoplasmic reticulum localized GFP signal (GFP_{er}) is available without heat shock (w/o HS) (a). In contrast, in leaf primordium 72 hours after heat shock (w/ HS), GFP_{er} is observed in continuous cell files (white arrowheads) along the mediolateral direction (b). (c) shows the magnification of a part of (b). (d-e) The distribution of inner cell division angle which is against the mediolateral axis labeled as blue (d) in a collection of optical cross sections of one entire tomato P3 from the tip to the base (e). (f, g) The distribution of epidermal cell division angle which is against the corresponding tangent labeled as blue (f) in a collection of optical cross sections of one entire tomato P3 from the tip to the base (g). Scale bars, 20 μ m in (a) and (b).

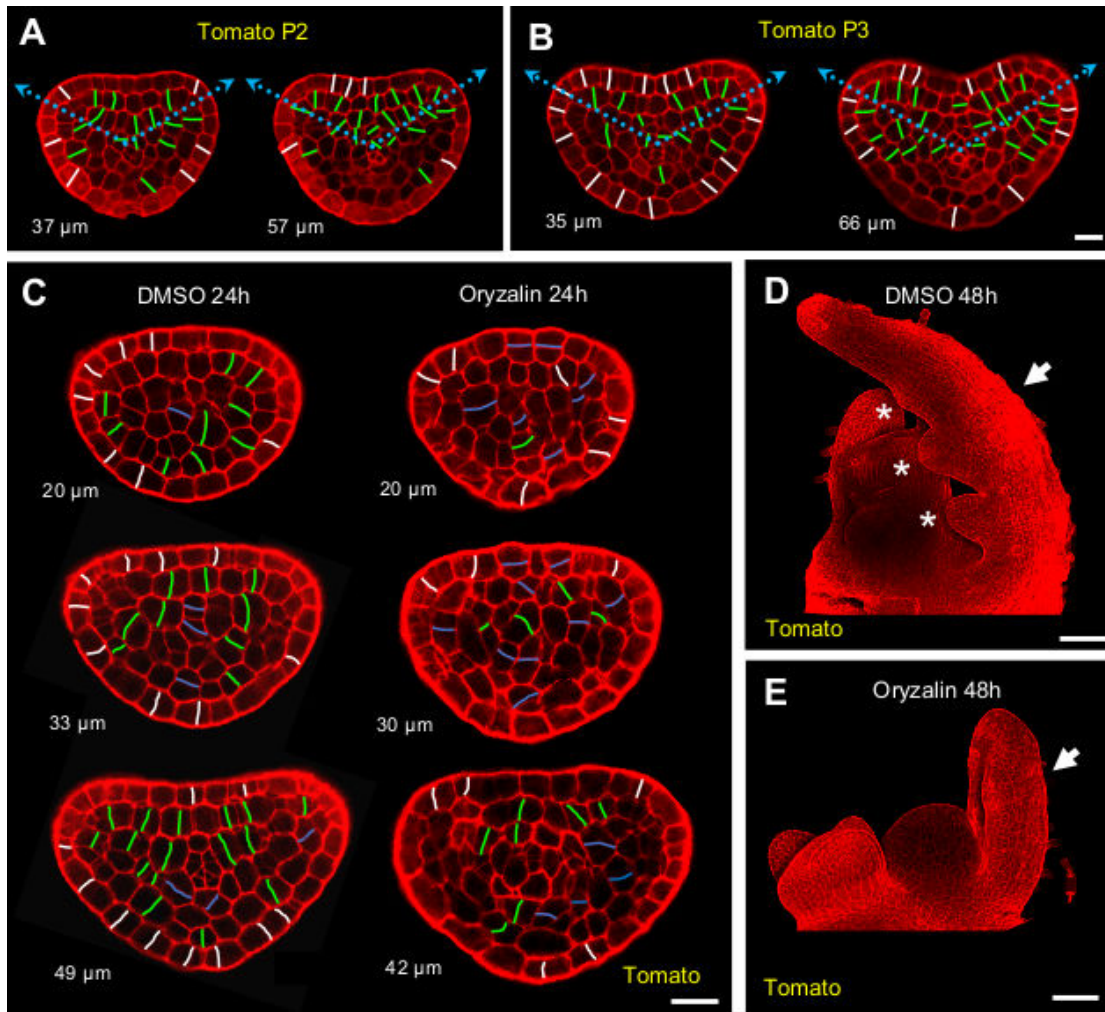


Figure 4.13: Importance of CMTs for orientated cell division and planar leaf formation. (a, b) Cell division pattern by mPS-PI staining in optical cross sections of tomato P2 (a) and P3 (b) show that the inner cells tend to divide perpendicularly to the mediolateral axis (blue line), while the epidermal cells exclusively perform periclinal divisions with the division plane perpendicular to the outer surface. The newly formed cell walls are labeled green for inner cells and white for epidermal cells. The depth of the optical cross sections from the tip of corresponding leaf primordium is shown. (c) Cell division pattern by mPS-PI staining in optical cross sections of tomato P3 24h after the treatment of DMSO (Mock) (left column) and oryzalin (right column). The newly formed cell walls are labeled green for inner cells and white for epidermal cells. The depth of the optical cross sections from the tip of corresponding leaf primordium is shown. (d, e) The morphology of tomato P3 48h after the treatment of DMSO (d) or oryzalin (e). Arrows indicate the primordia treated with chemicals. Leaf primordium treated with DMSO can perform normal anisotropic growth and generate lateral leaflet primordia (asterisk). The anisotropic growth and planar leaf form is compromised in oryzalin treated samples. Scale bars, 20 μm in (a-c), 100 μm in (d, e).

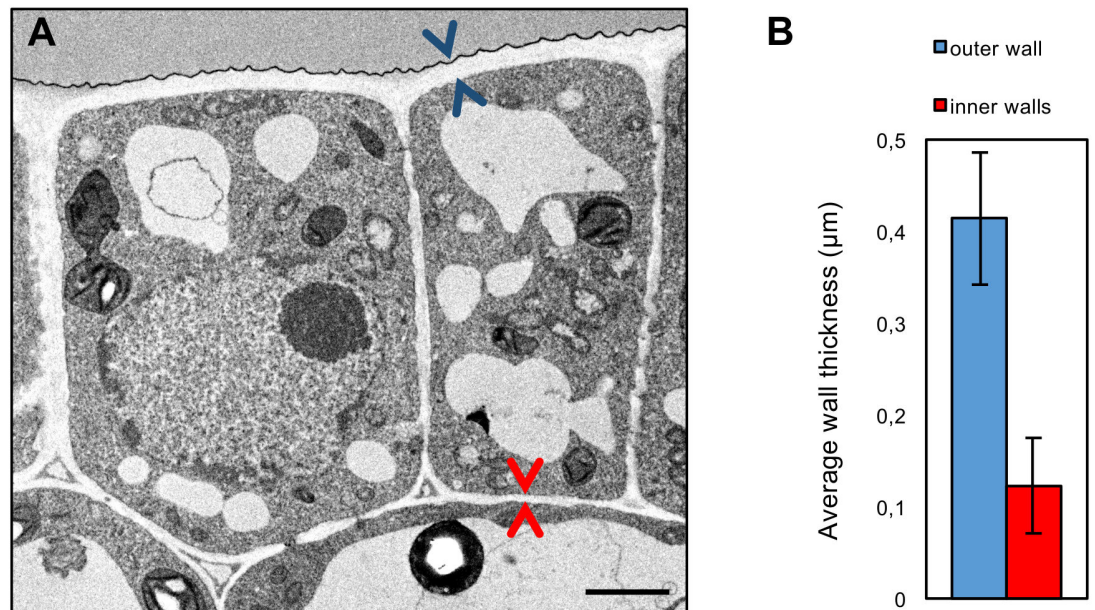


Figure 4.14: **Cell wall thickness in tomato leaf primordia.** (a) TEM micrograph of the epidermal cell lay in tomato P3 leaf primordium cross section showing the out cell wall (between blue arrowheads) is thicker than inner cell walls (between red arrowheads). (b) Quantification of (a). Scale bar, 2 μm in (a).

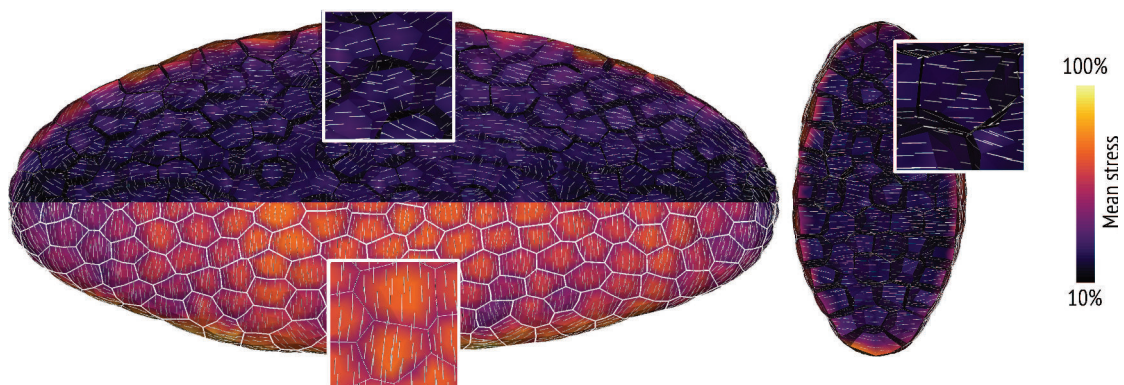


Figure 4.15: **Mechanical conflict generated by feedback on all walls.** When all walls show mechanical feedback, the outer layer will mainly orient its CMTs perpendicular to the longest axis of the ellipsoid and cause the structure to lengthen. This will cause resistance of the inner cells, which will orient their CMT along that axis. This will generate distortions in the long run. Note that in the axial plane, all CMTs orient anticlinally as also observed *in vivo*. Green arrows indicate main CMT orientations in different planes and layers.

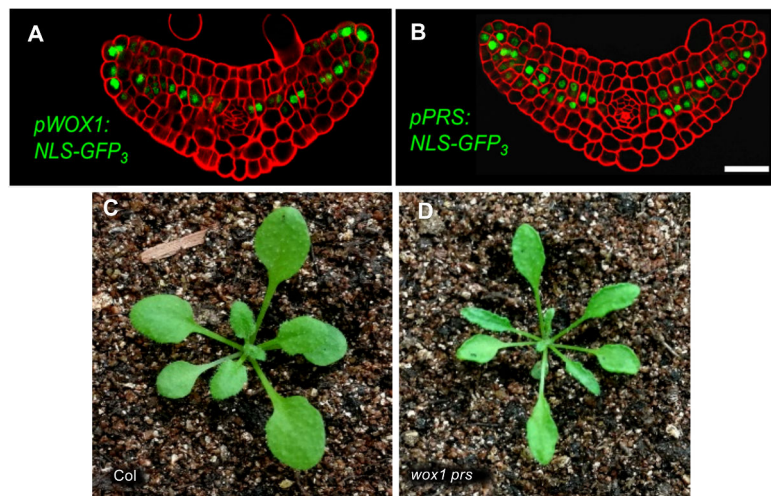


Figure 4.16: **Expression pattern of WOX genes and phenotype of wox1 prs in Arabidopsis.** (a, b) Expression patterns of pWOX1:NLS-GFP₃ (a) and pPRS:NLS-GFP₃ (b) in optical cross sections of Arabidopsis leaf primordia. Both reporters show GFP expression in the middle domain, while pWOX1 also shows additional activity in several cell layers in leaf margins. (c, d) Mature leaf phenotypes of 17 DAG old Col (c) and wox1 prs (d). Scale bar, 20 μm.

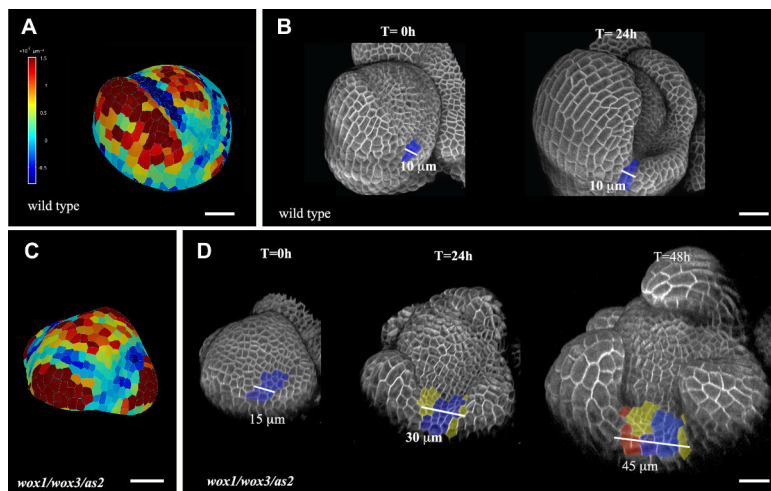


Figure 4.17: **Boundary formation in wild-type and wox1 prs as2 mutant flowers.** (a) Segmented 3D reconstruction of wild type (35S: GFP-Lti6b) at 0h, showing degree of Gaussian curvature. Blue color indicates negative curvature which is a marker for the boundary. (b) Confocal, 3D reconstruction showing the same flower bud at two time points. The site of negative curvature is marked blue. The width of this boundary does not change and remains about 10 μm wide. (c) Segmented 3D reconstruction of wox1 prs as2 (35S: GFP-Lti6b) at 0h. (d) Development of boundary in the triple mutant at three time points. The initial zone of negative curvature is slightly broader than the wild type and then gradually increase in size (from about 15 μm to 45 μm, marked with color from blue to yellow and red). Scale bars, 20 μm.

Chapter 5

Discussion

5.1 Towards more realistic models of the cell wall

The cell wall composition and structure can vary more or less rapidly, which confers cells the ability to modulate their growth in rate and direction, in an efficient manner. To model the 2D elastic properties of the wall, many models, like those presented in [Bassel et al. \(2014\)](#), [Boudon et al. \(2015\)](#), [Bozorg et al. \(2014, 2016\)](#), have considered simple orthotropic¹ materials, typically in the form of Saint-Venant Kirchhoff-like strain-stress relationships, the parameters of which did not reflect the composite structure of the wall in any detail.

The elastic model developed in this thesis is an effort to connect the wall elastic parameters to the mesoscopic structure of the wall, focusing on CMFs. This was done by representing CMFs as short linear springs under affine deformation, adapting a model originally designed to represent the elasticity of nonliving fibrous materials like paper ([Cox, 1952](#)). A similar approach was developed in [Armezzani et al. \(2018\)](#) (for the SAM epidermis), also based on a Cox-like description of the stiffness tensor. Yet, this model did not represent the dynamical properties of CMFs and CMTs. Here, by coupling the stiffness model with a kinetic equation governing the polymerization/depolymerization of CMFs, we derived a dynamic constitutive behavior of the wall, taking into account stress feedback and growth-induced advection, in a physically motivated manner.

In this model, we assumed that CMTs were stabilized in the direction of main stress, which in average results in a directional selection, promoting CMT coalignment in the

¹Structurally symmetric around the axes of anisotropy.

axis of main stress (Eq. (2.9)). To represent the dynamic modification of the wall elasticity, we expressed the deposition of CMFs by transmembrane cellulose synthases as a function of CMTs distribution. One of the main assets of our approach, compared to other published works, is that the dynamic evolution of the wall elasticity explicitly depends on the kinetic constants of the chemical processes involved in wall turnover. In addition, coupling chemical kinetics with a growth-induced advection of the fiber raises a number of questions (Chapter 3). In particular, we discussed in Chapter 3 some minimal conditions on cellulose kinetic processes, allowing for a cell to maintain a homeostatic stiffness, while keeping the ability to perform a rapid change in growth direction if required.

As a limit of our approach, the affine deformation hypothesis (which is a prerequisite for the constitutive behavior developed in Chapters 2 and 3) will by construction not capture the possible heterogeneities of strain that may appear within the wall at a small scale. These could emerge when the dynamic evolution of the wall structure occurs in a time scale comparable to that of growth, resulting in differences in strain between the wall constituents. These may confer interesting properties to the mechanics of the wall, as for example modeled by Dyson and colleagues in the context of hemicellulose synthesis and breakage (Dyson et al., 2012). One may also remark that simple, linear strain-stress constitutive relations (like that used in this thesis), may also lack realism. In fact, more nonlinear elastic effects are frequent in biological mechanical systems (Chagnon et al., 2015). These effects are not fully characterized in the context of the meristem, which is not easily accessible to mechanical testing, although Kierzkowski *et al.* have recently proposed, on an experimental ground, that nonlinear effects may occur at the SAM, in the form of a strain hardening effect (Kierzkowski et al., 2012).

This thesis is an effort to describe the link between wall structure and wall elasticity. Yet, the plastic regime was described in a phenomenological manner, inspired by Lockhart's approach (Lockhart, 1965), which is itself an empirical rheological formulation. Finer growth laws were proposed to reflect the dynamics of wall turnover and irreversible creep. For example, Dyson and colleagues established a mechanistic explanation of Lockhart's law, where the viscous wall creep is the result of an imbalance between fast wall expansion (causing hemicellulose breakage) and slow wall turnover (Dyson et al., 2012). This model was later adapted to 2D growth *in silico* in Fozard et al. (2013).

5.2 Stress feedback in multi-layered tissues

Often enough, theoretical and experimental studies on meristem mechanics have focused either on the outermost cell layer or on the outer cell wall. In fact, according to the prevailing hypothesis, these bear most part of the tissue stresses (at least in the SAM, see [Beauzamy et al., 2015](#), and references therein). This allows to assimilate the whole tissue to a pressurized shell, neglecting the inner walls of the tissue (as in Chapters 2 and 3 and [Bozorg et al., 2014, 2016](#), [Hamant et al., 2008](#), [Kierzkowski et al., 2012](#), for example).

The actual mechanical role of these inner walls is yet not straightforward. For instance, it was suggested, by simulation, that the presence of inner walls can yield stress patterns different from those predicted through a pressurized-shell hypothesis ([Ali et al., 2019](#)). We also stressed that most of the shell-based models, coupling growth and stress feedback, in the literature ([Bozorg et al., 2014](#), [Hamant et al., 2008](#)) had been used in the context of prolate axisymmetric organs (typically stems). In modeling flat structures like leaves (which have a different type of symmetry), it appeared that a shell-based approach is probably not appropriate to model planar expansion (Chapters 3 and 4). To represent the latter, we needed a finer representation of the tissue, taking the inner walls into account (Chapter 4). In fact, we suggest that oriented CMFs at inner walls are crucial for this type of growth regime. This would mean that a shell-based approximation is not only coarse, but fundamentally insufficient to represent planar expansion.

To model the expansion of this kind of extremely asymmetric structures, growth and stress feedback were simulated using ellipsoidal meshes, containing both outer periclinal wall and inner tissues, and composed of 800 cells. To generate these meshes, a dedicated method (presented in Chapter 4) was implemented, based on [Cerutti et al. \(2017\)](#).

This full 3D approach allowed us to predict fine properties emerging from the coupling of growth and stress feedback in 3D alveolar tissues. We first observed that, *in silico*, a complete “activation” of the feedback in all membranes of the tissues, over-constrained the system, which resulted in a loss of cohesion in orientation between inner and outer CMTs/CMFs (see Chapter 4 and Fig. 4.15). This can be explained by the anisotropy of the outer wall that promotes apico-basal growth, which in turn stretches the inner cells. These dynamically respond to this traction by orienting their CMTs accordingly. The anisotropy of the surface walls was not consistently observed *in vivo*, as in leaves and sepals, the outer periclinal microtubules rapidly became disorganized. If our modeling assumptions are correct, in particular if a stress feedback does exist in leaves, this could indicate that a fine regulation of the system’s sensitivity is required, preventing

the effect predicted by simulation to occur in reality. Disabling the stress feedback on the outer wall of the tissues in our simulations (which is consistent with *in vivo* observations) in fact further allowed to generate more stable, flat growing shapes. This potentially provides a mechanistic explanation of anisotropic leaf growth.

5.3 Cell division and meshes

Another crucial aspect of development is cell division. This aspect was not addressed in this thesis. While simulating division in discrete 2D approaches (e.g. spring-based models) is trivial, no satisfactory computational solution has so far been proposed to perform this operation in cell-scale 3D models like those used in Chapter 4. In fact, simulating a cell division means adding a new wall within a cell and arbitrarily cutting the already existing walls in two. Performing this cut directly on the finite elements forming a given cell wall would probably result in a dramatic decrease of mesh quality. Due to this limitation, the 3D simulations of growth presented in Chapter 4 do not integrate division and, for this reason, can only capture short developmental times.

Correlatively, anisotropic growth results in distortions of meshes that cause the FEM to become increasingly unreliable in the long run. A major future challenge, which can be a subject of research *per se*, is to design algorithms to dynamically update meshes and simulate cell division in an efficient manner. The aim here would be to sustain sufficient mesh quality in time, while ensuring the continuity of the various properties associated with the mesh triangles (in particular the strain). Remeshing/division methods tailored for planar (Fozard et al., 2013) or 2D curved (Bozorg et al., 2016) cells can be of interest while addressing those various issues with 3D polyhedral cells.

5.4 Do plant cells “measure” stress?

More than a decade ago, Hamant and coworkers reexamined the seminal idea (originally proposed by Green and King, 1966, in *Nitella*) of a causal relationship between mechanics and tissue physiology, mediated *via* microtubules (Hamant et al., 2008). The appealing idea that forces may provide a positional and directional cue to cells has become increasingly popular in the plant biology literature and has motivated various experimental (e.g. Burian et al., 2013, Hervieux et al., 2016, 2017, Sampathkumar et al., 2014a, Sapala et al., 2018, Uyttewaal et al., 2012) or theoretical (Alim et al., 2012, Bozorg et al., 2014, Sapala et al., 2018) investigations. To date however, the hypothesis

of a mechanical control of plant morphogenesis still lacks a clear mechanistic understanding (for reviews, see [Hamant and Haswell, 2017](#), [Hamant et al., 2019](#), [Landrein and Hamant, 2013a](#)).

Despite the increasing experimental evidence underpinning the stress feedback hypothesis ([Hamant et al., 2019](#)), one may object that the original terminology employed in [Hamant et al. \(2008\)](#) *inter alia*, proposing that “stresses orient microtubules” remains unsatisfactory from a physical standpoint. In fact, defining a *force* as a physical object on its own, other than the *physical cause of acceleration*, is not self-evident (see [Feynman et al., 1965](#), chap. 12). Likewise, it is known that a force cannot be measured by other means than its effect (motion). That cells actively react to *forces* (and *stresses*) is hence a rather abstract statement, that does not easily allow to picture a reasonable physical and chemical mechanism. Indeed, evidence wants that the “perception” of a force should in any way rely upon the perception of a *strain*.

Could that strain be the tangential elastic strain of the walls? In anisotropic materials like the plant cell wall, the respective main axes of stress and strain may not coincide. Instead they are perpendicular in a number of cases, as explored in Chapter 2. Thereby, it is equally reasonable to suggest that microtubules align *perpendicular* to main strain, rather than along main stress ([Schopfer, 2009](#)). Nonetheless, this scenario fails in the rare cases where main stress and strain coincide, like in the concave crease around organs, where CMTs, main strain and main stress are supposed to be parallel ([Ali et al., 2019](#), [Burian et al., 2013](#), [Dumais and Kwiatkowska, 2002](#)). Theoretical modeling also tends to disprove such a possibility, by suggesting that an alignment of CMTs/CMFs perpendicular to strain would be unstable ([Bozorg et al., 2014](#)).

One could also imagine that CMTs could be anchored to the wall and put under tension by wall deformation. As suggested by [Inoue et al. \(2016\)](#) for example, CMTs under tension would in turn stabilize (as also discussed in [Hamant et al., 2019](#)). However, according to this scenario, CMTs would always align with the wall main strain. This would not be compatible with cases where main stress and strain are perpendicular and where CMTs are aligned along the main stress direction. Alternatively, CMT organization could be controlled by means of other intermediate molecular actors. CLASP proteins for instance, that accumulate at cell edges, regulate CMT alignment, by promoting CMT stability ([Ambrose et al., 2011](#)). A correlation between predicted stress and CLASP – or any other chemical actor – distribution, which has not yet been explored to my knowledge, could potentially provide insight into the problem.

The scenario of an indirect chemical regulation of CMT alignment would require that some information on wall stress would be chemically transferred to CMTs. This could be based on the tension-induced stabilization of some molecular actor embedded within

the wall, acting as a stress sensor. However, how such a mechanism could provide a directional information is not self-evident. For example, this sensor could have an asymmetric – e.g. filamentous – shape, resulting, through stabilization, in an average coalignment, and therefore a directional information. Yet, again, this tension should *a priori* be caused by wall strain (perpendicular to the wall stress): we find ourselves in the same predicament as before.

In this thesis, I have mostly tried to represent the emergent effects of a stress feedback, without questioning its actual physical foundations. It yet appeared to me that the various explanatory scenarios proposed in the literature are not completely satisfactory and are often abstract. In this conclusion, I would like to sketch a possible general mechanism, based on parsimonious physical principles and inspired by the mechanoperception mechanisms better identified in animals (Charras and Yap, 2018). Briefly, the idea developed here is that the strain energy stored nonuniformly by the various adhesion zones around the cell, has the potential to provide an anisotropic, cell-scale chemical information on the external loading.

In the majority of plant cellular biomechanical models, it is more or less implicitly assumed that stresses are borne by the cell walls only, as cells are forced to be in perfect contact. This was for example an assumption of our model, where we assumed that two cells shared the same wall. In reality, each cell is embedded within its own wall exoskeleton, and contiguous cells are connected to each other through a sticky, pectin-rich *middle lamella* (Daher and Braybrook, 2015, Jarvis et al., 2003, Knox, 1992). This middle lamella potentially plays an important role in transferring mechanical forces from cell to cell, as experimentally investigated by Verger and coworkers, on *Arabidopsis* hypocotyl and cotyledon (Verger et al., 2018).

Consider a cell subject to an anisotropic – say tensile – external loading. *A priori*, the strain energy stored in the middle lamella should be heterogeneously distributed around this cell. In fact, zones of highest strain energy density should be those where cells tend to be split apart with the strongest pull, *i.e.* where the lamella is the most mechanically solicited. In a multicellular context the main axis of tissue-scale stress could be normal to these zones. This was suggested by Verger et al. who evidenced that in the adhesion-deficient mutant *quasimodo1* (Bouton et al., 2002), cells detach along the lines of maximal predicted tension (Verger et al., 2018).

The heterogeneous distribution of adhesion strain energy provides a cell-scale directional information on the stress field. Mechanistically, this information could be transduced through a cascade of chemical reactions (that would remain to be characterized), from an adhesion strain sensor, all the way to CMT polymerization/depolymerization processes. CMT nucleation or stabilization could be for example promoted by specific

chemical actors nonuniformly distributed around a cell (Fig. 5.1). In addition, CMT behavior is known to depend on the mutual interactions between CMTs (Dixit and Cyr, 2004), which may amplify their coalignment.

This scenario, summarized in Fig. 5.1 could provide a conceptual basis for a mechanistic explanation of the stress feedback. In animals, adhesion-based mechanosensing – mediated *via* cadherins for instance – was better characterized (see Charras and Yap, 2018). In plants, the above-detailed mechanism could potentially account for the alignment of periclinal CMTs. Yet, it remains unclear whether it will be the case for other membranes. For instance, anticlinal membranes at the SAM often display CMTs pointing towards the outside, irrespective of how anisotropically distributed periclinal CMTs are. Further knowledge on the mechanical state of the middle lamella between the epidermis and the second cell layer is required. It also appears that this mechanism could provide a cell-scale information. Whether it can account for subcellular CMT patterns like those reported in Sampathkumar et al. (2014a) is unclear.

To my knowledge, this general scenario has not yet been explored, neither experimentally nor theoretically. Modeling could provide a tool to assess the theoretical plausibility of such a mechanism, from a macroscopic standpoint, for example by first allowing for the quantitative estimation of the adhesion-related strain energy distribution.

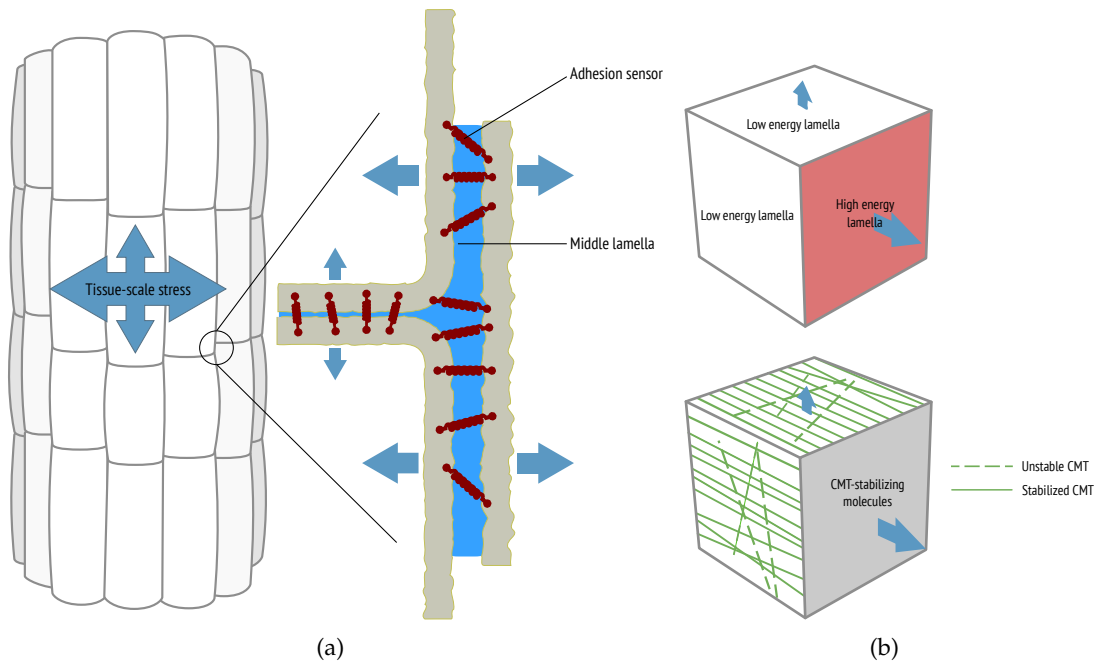


Figure 5.1: Sketch of a conceptual adhesion-based mechanism for the perception of tissue-scale stresses in plants. A cell subject to an anisotropic loading would store adhesion-related strain energy in a differential manner, depending on the orientation of the membranes with respect to loading axis. In turn, the various levels of strain energy stored in the middle lamella may provide a directional information allowing to regulate CMT alignment. (a) schematic cylindrical organ (stress is mainly circumferential), close-up on the middle lamella containing mechanical sensors. (b) Cartoon of a cell aligning its microtubules along the axis of main tensile loading. The strain energy stored in the middle lamella is non uniformly distributed around the cell. This in turn modulates some chemical process (for example stabilization of some molecular actor) that provides the directional information.

Glossary

apoptosis	Programmed cell death. 7
Cauchy stress	Physical quantity that expresses the internal forces acting within an infinitesimal volume of matter in the current configuration. ... 39
cell division	Process by which a parent cell divides into two daughter cells. 7
cell growth	Irreversible expansion of the cells due to turgor-induced forces. 6
cell wall 6
CMFs	cellulose microfibrils..... 24
CMTs	cortical microtubules 25
CSCs	cellulose synthase complexes 25
deformation gradient 9
elasticity tensor 23, 36
Eulerian strain rate tensor 8
FEM	finite element method 15

Green-Lagrangian strain	Geometric measure of how much a displacement locally differs from a rigid body displacement. 22, 23
growth tensor 10
meristem 5
morphoelasticity 2
morphogenesis 1
multiplicative decomposition 10
organogenesis 5
osmosis	Spontaneous movement of water through the (semipermeable) cell membranes due to a difference in solute concentration between both sides of the membrane. This allows cells to absorb water. 6
rate of growth 10
SAM	shoot apical meristem. 5
second Piola-Kirchhoff stress	Physical quantity that expresses the internal forces within an infinitesimal volume of matter in the reference configuration. 23
spin tensor 8
strain energy density	Volumetric density of potential strain energy (in J.m^{-3}) which is stored by a mechanical system undergoing a deformation. 10

stress	Physical quantity that expresses the internal force within an infinitesimal volume of matter. It can be mathematically expressed in various manners, in particular depending on whether we express the forces in the current or reference configuration of the material. 10
tissue	A contiguous set of cells..... 2
turgor pressure	Cell internal hydrostatic pressure resulting from osmotic water intake, which is resisted by the cell wall. 2, 6
velocity gradient 8

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